

Verification And Validation Of TMAP7

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ABSTRACT

The Tritium Migration Analysis Program, Version 7 (TMAP7) code is an update of TMAP4, an earlier version that was verified and validated in support of the International Thermonuclear Experimental Reactor (ITER) program and of the intermediate version TMAP2000. It has undergone several revisions. The current one includes radioactive decay, multiple trap capability, more realistic treatment of heteronuclear molecular formation at surfaces, processes that involve surface-only species, and a number of other improvements. Prior to code utilization, it needed to be verified and validated to ensure that the code is performing as it was intended and that its predictions are consistent with physical reality. To that end, the demonstration and comparison problems cited here show that the code results agree with analytical solutions for select problems where analytical solutions are straightforward or with results from other verified and validated codes, and that actual experimental results can be accurately replicated using reasonable models with this code. These results and their documentation in this report are necessary steps in the qualification of TMAP7 for its intended service. This revision updates results using TMAP7.1, which corrected some code deficiencies found in TMAP7.

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1.0 OVERVIEW

The TMAP Code was written at the Idaho National Engineering and Environmental Laboratory by Brad Merrill and James Jones in the late 1980s as a tool for safety analysis of systems involving tritium. ¹Since then it has been upgraded to TMAP4 and has been used in numerous applications including experiments supporting fusion safety, predictions for advanced systems such as the International Thermonuclear Experimental Reactor (ITER), and estimates involving tritium production technologies. The code's further upgrade to TMAP2000² and then to TMAP7 was accomplished in response to several needs. TMAP and TMAP4 had the capacity to deal with only a single trap for diffusing gaseous species in solid structures. TMAP7 includes up to three separate traps and up to 10 diffusing species. The original code had difficulty dealing with heteronuclear molecule formation such as HD. That has been removed. Under solution-law dependent diffusion boundary conditions, such as Sieverts' law, TMAP7 automatically generates heteronuclear molecular partial pressures and surface flows when solubilities and partial pressures of the homonuclear molecular species are provided. A further sophistication is the addition of non-diffusing surface species. Atoms such as oxygen or nitrogen or complexes such as hydroxyl radicals on metal surfaces are sometimes important in molecule formation with diffusing hydrogen isotopes but do not themselves diffuse appreciably in the material. TMAP7 will accommodate up to 30 such surface species, allowing the user to specify relationships between those surface concentrations and partial pressures of gaseous species above the surfaces or to form them dynamically by combining diffusion species or other surface species. Additionally, TMAP7 allows the user to include a surface binding energy and an adsorption barrier energy and includes asymmetrical diffusion between the surface sites and regular diffusion sites in the bulk. All of the previously existing features for heat transfer, flows between enclosures, and chemical reactions within the enclosures have been retained, but the allowed problem size and complexity have been increased to take advantage of the greater memory and speed available on modern computers. One feature unique to TMAP7 is the addition of radioactive decay for both trapped and mobile species. Another is the ability to initialize distributed parameters such as initial mobile atom, trapped atom, or trap concentrations using selected mathematical functions. Also, time-dependent temperatures and pressures can be specified in *boundary* enclosures and for surface concentrations of diffusion species.

Since its release in November 2004, TMAP7 was found to have a few minor errors that prevented correct results from being obtained in certain problems. Those errors have been corrected in TMAP7.1, which will not be specifically referred to hereafter except as TMAP7. Results included in this revision are for the TMAP7.1 version of the code.

The verification and validation process normally involves two steps. The verification process is a careful examination of the code to ensure that the coding faithfully reproduces the mathematical model and that the code is well written and efficient. That process was pursued extensively with TMAP4 but has not been done independently of code development here. The basic architecture of the code remains the same, although a number of minor changes were required to work with the GNU FORTRAN 77, selected for distribution with the code. There are also new components and a few new subroutines. These have been carefully evaluated for coding accuracy, but the demonstration of their success is in the high fidelity the code provides to the sample problems. Those sample problems constitute the validation of the code and provide the basis for what is presented here.

There are two main sections to this report. The first exercises TMAP7 in each of its major capability areas using specialized problems, showing that the results computed by TMAP7 are in good agreement with “known” results. This demonstrates that the code’s functional tools are performing properly. The second part of the report provides a comparison of TMAP7 results with experimental results to show the general utility of the code in modeling reality.

2.0 SPECIALIZED PROBLEMS

Computational capabilities of TMAP7 lie in six major areas: diffusion and trapping within structures and surface processes, heat transfer, chemical reactions in enclosures, bulk fluid flows, chemical equilibrium and radioactive decay. The demonstration problems that follow are grouped into those areas.

Problems 1a-1e exercise TMAP7’s mass transfer capabilities

Problems 1f (a-c) demonstrate TMAP7's heat transfer functions

Problems 1g (a-c) model enclosure reactions

Problems 1h (a-b) deal with enclosure flow

Problem 1i (a-b) verify chemical reactions in enclosures and on surfaces are correct

Problem 1j demonstrates radioactive decay.

The descriptions of these problems include a statement of the problem, a description of the modeling used in setting up the problem for TMAP7, and a comparison of the TMAP7 results with “known” solutions from literature or other sources. Appendix A is the derivation for the surface equilibrium model used in problem 1i (b). Appendix B contains the input code listings for each of the problems cited in the report.

The file names assigned to the various problems appear in parentheses in the headings for the problem descriptions. Input files carry the *.inp* extension, output or *codeout* files have *.out*, and plot data files (*pltdata*) terminate with the *.plt* extension.

Theoretical results were calculated using Microsoft Excel™, and TMAP7 calculations were obtained in two working environments. One used Windows XP™ on a Dell Optiplex GX 260 and on a custom-built Pentium 4 3.0-GHz machine. The other was Windows ME™ running on a Dell Dimension XPS R450 and on a Dell Latitude C-600 laptop computer.

2.1 Problem 1a: Diffusion from a Depleting Source (Val-1a)

This diffusion problem models an enclosure that is pre-charged with a fixed quantity of tritium. At time $t > 0$, the tritium is allowed to diffuse through a finite slab of SiC, initially at zero concentration. The surface of the slab in contact with the source is assumed to be in equilibrium with the source enclosure. The boundary condition at the exit side of the slab is kept constant at zero concentration for all time. The concentration of the enclosure is then calculated for different times and reported as a fractional release. There are no trapping effects active in the slab.

Carslaw and Jaeger³ give the analytical solution for an analogous heat transfer problem from which the solute concentration profile in the membrane is

$$C(x, t) = 2SP_0L \sum_{n=1}^{\infty} \frac{\exp(-\alpha_n^2 Dt) \sin(\alpha_n x)}{[l(\alpha_n^2 + L^2) + L] \sin(\alpha_n l)} \quad (1)$$

where

$$\alpha_n = \frac{L}{\tan(\alpha_n l)} \quad (2)$$

$$L = \frac{STAk}{V} \quad (3)$$

Here

A = cross-sectional area of the slab ($2.16 \times 10^{-6} \text{ m}^2$)

D = diffusivity of tritium (SiC assumed: $2.62238\text{E-}11 \text{ m}^2/\text{s}$ at 2373 K)

k = Boltzmann's constant ($1.38065 \times 10^{-23} \text{ J/K}$)

l = thickness of the slab ($3.30 \times 10^{-5} \text{ m}$)

S = solubility of tritium (SiC assumed: $7.244\text{E}22/T \text{ atom/m}^3/\text{Pa}$)

T = temperature (2373 K)

V = volume of the enclosure ($5.20 \times 10^{-11} \text{ m}^3$)

We apply Henry's law to the concentration at $x = l$ to find the gas pressure in the enclosure

$$P(t) = \frac{C(l, t)}{S} = 2P_0L \sum_{n=1}^{\infty} \frac{\exp(-\alpha_n^2 Dt)}{l(\alpha_n^2 + L^2) + L} \quad (4)$$

and finally the release fraction

$$FR = 1 - \frac{P(t)}{P_0} = 1 - 2L \sum_{n=1}^{\infty} \frac{\exp(-\alpha_n^2 Dt)}{l(\alpha_n^2 + L^2) + L} \quad (5)$$

Some of the values obtained from Equation (5) and from TMAP7 are compared in Table 1. Ten terms were included in the sum of (5) so that even at $t = 1 \text{ s}$, the last term was less than 10^{-10} of the sum. The variance between the analytical solution and the computed solution from TMAP7 is defined by Equation (6)

$$Variance = \frac{TMAP7 - Analytical}{Analytical} \quad (6)$$

Table 1. Fractional release of tritium from depleting source problem Val-1a.

Time (s)	TMAP7	Theory	Variance
0	0.00000	0.00000	0.00000
1	0.19383	0.20439	0.05169
2	0.26151	0.26640	0.01837
3	0.30708	0.31022	0.01011
4	0.34181	0.34409	0.00662
5	0.36998	0.37172	0.00467

Time (s)	TMAP7	Theory	Variance
6	0.39375	0.39510	0.00342
7	0.41434	0.41541	0.00257
8	0.43260	0.43343	0.00191
9	0.44906	0.44971	0.00144
10	0.46415	0.46464	0.00105
11	0.47815	0.47851	0.00076
12	0.49127	0.49153	0.00054
13	0.50368	0.50386	0.00036
14	0.51550	0.51562	0.00023
15	0.52683	0.52690	0.00013
16	0.53774	0.53777	0.00006
17	0.54827	0.54827	0.00001
18	0.55848	0.55845	-0.00005
19	0.56839	0.56834	-0.00008
20	0.57802	0.57797	-0.00010
21	0.58740	0.58734	-0.00011
22	0.59655	0.59648	-0.00012
23	0.60547	0.60540	-0.00012
24	0.61419	0.61411	-0.00013
25	0.62270	0.62261	-0.00013

The variance decreases almost monotonically for $t > 25$ s. Figure 1 shows the comparison for the first 140 s.

A further comparison may be made by noting that the surface flux at $x = 0$ is

$$J = D \frac{\partial C(x, t)}{\partial x} \bigg|_{x=0} = 2SP_0LD \sum_{n=1}^{\infty} \frac{\exp(-\alpha_n^2 Dt) \alpha_n}{[l(\alpha_n^2 + L^2) + L] \sin(\alpha_n l)} \quad (7)$$

A comparison of results for flux through the free surface is shown in Figure 2.

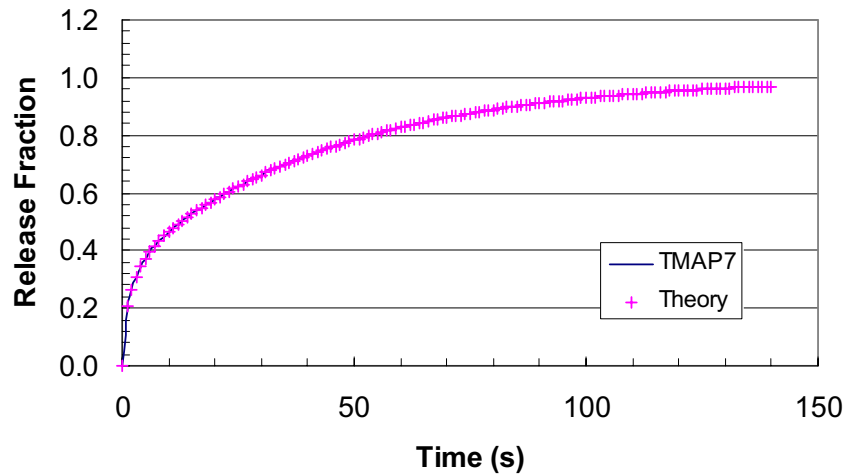


Figure 1. Fractional release of tritium from an enclosure through SiC in depleting source demonstration problem (Val-1a).

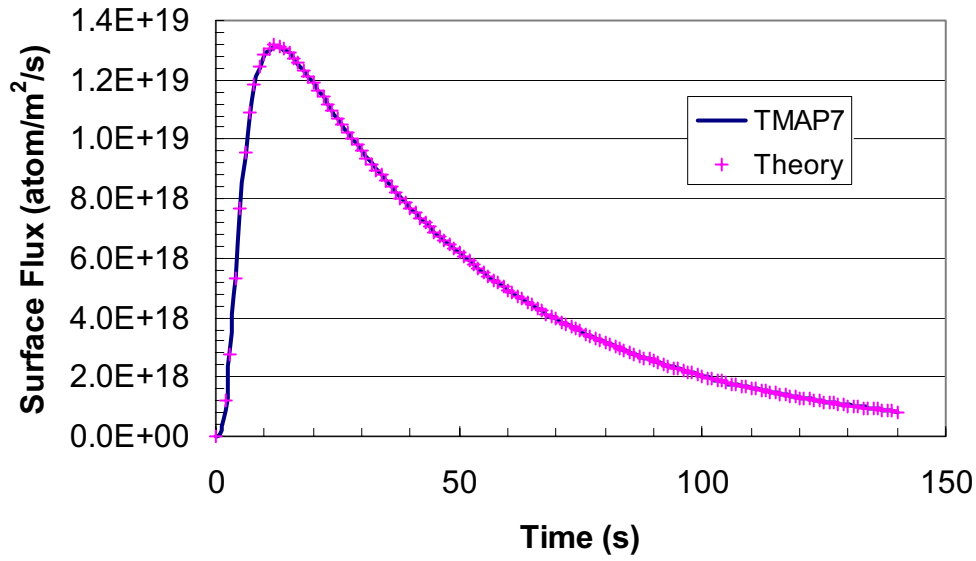


Figure 2. Atom flux through outside face of membrane for depleting source problem (Val-1a).

2.2 Problem 1b: Diffusion in a Semi-Infinite Slab with Constant-Source Boundary (Val-1b)

This model is designed to test the basic Fick's-law diffusion. A semi-infinite slab is defined with a constant concentration boundary condition. The initial concentration of the slab is zero for time, $t \leq 0$ seconds. At time $t > 0$, the diffusion is allowed to proceed. The slab is assumed to have no traps. Three comparisons are shown; a transient concentration history at a given location, a spatial concentration profile at a given time, and the variation of flux into the slab surface. These are compared with analytical results.

Carslaw and Jaeger⁴ give the analytical solution to the time-dependent concentration profile as

$$C(x,t) = C_o \operatorname{erfc}\left(\frac{x}{2\sqrt{Dt}}\right). \quad (8)$$

where

$C(x,t)$ = diffusion species concentration at position x and time t

C_o = concentration of the diffusing species at the free surface (1.0 atoms/m³)

D = diffusivity (1.0 m²/s).

The solution of Equation (8) was found using Microsoft Excel using the series expansion given in CRC Standard Mathematical Tables and Formulae⁵. This expansion is

$$\operatorname{erfc}(x) = 1 - \operatorname{erf}(x) = 1 - \frac{2}{\sqrt{\pi}} \left(x - \frac{x^3}{3} + \frac{1}{2!} \frac{x^5}{5} - \frac{1}{3!} \frac{x^7}{7} + \frac{1}{4!} \frac{x^9}{9} \dots \right). \quad (9)$$

Enough terms were taken in this expansion to ensure that the last term contributed less than 1.0×10^{-12} for all values of the argument.

Two comparisons were made for this model between the values of Equation (8) and results from TMAP7. The first comparison was made for times ranging from $t = 0$ to 30 s at a distance from the surface of $x = 0.15$ m. The disagreement between Equation (8) and TMAP7 was less than 0.05% at $t = 1$ sec. The variance decreased with time, declining quickly to 0.001%. These values are listed in Table 2.

Table 2. Concentration profile at $x = 0.15$ m for problem Val-1b, diffusion in a semi-infinite slab.

Time (s)	TMAP7	Theory	Variation 1
0	0.000000	0.000000	0.000000
1	0.915140	0.915530	-0.000426
2	0.940080	0.940215	-0.000143
3	0.951100	0.951170	-0.000074
4	0.957660	0.957706	-0.000048
5	0.962130	0.962167	-0.000039
6	0.965430	0.965461	-0.000033
7	0.968000	0.968022	-0.000023
8	0.970070	0.970086	-0.000017
9	0.971780	0.971796	-0.000017
10	0.973230	0.973243	-0.000014
11	0.974480	0.974488	-0.000008
12	0.975560	0.975574	-0.000014
13	0.976520	0.976532	-0.000012
14	0.977380	0.977385	-0.000005
15	0.978150	0.978152	-0.000002
16	0.978840	0.978845	-0.000005
17	0.979470	0.979477	-0.000007
18	0.980050	0.980055	-0.000005
19	0.980580	0.980587	-0.000007
20	0.981070	0.981078	-0.000008
21	0.981530	0.981534	-0.000004
22	0.981950	0.981959	-0.000009
23	0.982350	0.982355	-0.000005
24	0.982720	0.982727	-0.000007
25	0.983070	0.983076	-0.000006
26	0.983400	0.983404	-0.000004
27	0.983710	0.983714	-0.000004
28	0.984010	0.984008	0.000002
29	0.984280	0.984286	-0.000006
30	0.984550	0.984550	0.000000

The second comparison examined the concentration profile from $x = 0.05$ to 19.8 m at increments of 0.1 m at time, $t = 25$ s. The variance between Equation (8) and TMAP7 is small, exceeding 0.1% only at depths greater than 6 m. The comparison of these values can be seen in Table 3, listing values to 11.9 m, and in Figure 3, out to 19.7 m.

Table 3. Concentration profile (atom/m³) at $t = 25$ sec for diffusion in a semi-infinite slab.

x (m)	TMAP7	Theory	Variance
0	1.00E-00	1.000000	0.000000
0.5	9.38E-01	0.943628	-0.005964
1.1	8.71E-01	0.876377	-0.006364
1.7	8.05E-01	0.810008	-0.006775
2.3	7.40E-01	0.744977	-0.007191
2.9	6.77E-01	0.681717	-0.007623
3.5	6.16E-01	0.620618	-0.008053
4.1	5.57E-01	0.562031	-0.008506
4.7	5.02E-01	0.506255	-0.008958
5.3	4.49E-01	0.453536	-0.009406
5.9	4.00E-01	0.404064	-0.009883
6.5	3.54E-01	0.357971	-0.010366
7.1	3.12E-01	0.315334	-0.010828
7.7	2.73E-01	0.276178	-0.011327
8.3	2.38E-01	0.240476	-0.011836
8.9	2.06E-01	0.208157	-0.012331
9.5	1.77E-01	0.179109	-0.012893
10.1	1.51E-01	0.153190	-0.013444
10.7	1.28E-01	0.130227	-0.014104
11.3	1.08E-01	0.110029	-0.014899
11.9	9.09E-02	0.092392	-0.015767

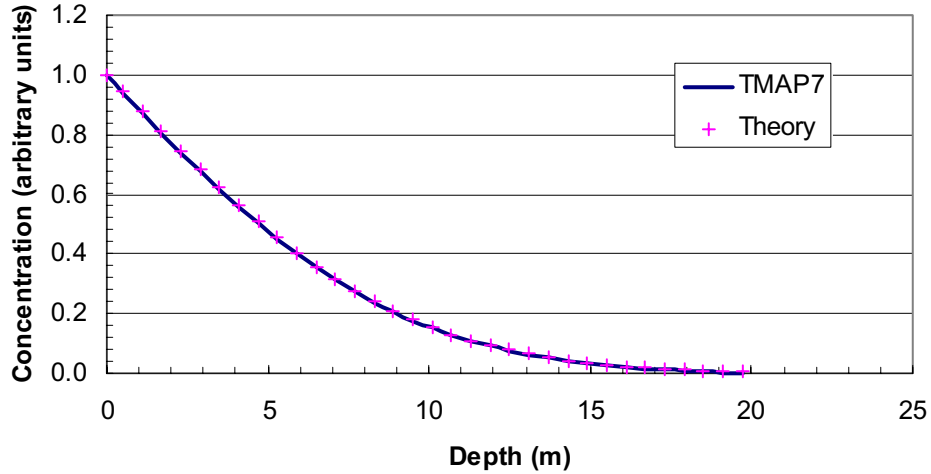


Figure 3. Concentration profile in a semi-infinite slab of SiC after 25 s from problem Val-1b.

The third, and final, comparison for this problem was the comparison of the diffusive flux into the slab. The flux into or out of a slab is proportional to the concentration gradient in the x direction at the slab surface. The solution⁶ is given by

$$J = C_o \sqrt{\frac{D}{t\pi}} \exp\left(\frac{x}{2\sqrt{Dt}}\right) \quad (10)$$

The values of Equation (10) were found using Microsoft Excel. They were compared to the values obtained from TMAP7 and can be seen in Table 4. The variance is never greater than 0.44%.

Table 4. Flux (atom/m² sec) into semi-infinite slab from a constant source

Time (s)	TMAP7	Theory	Variance
0	1.000000	1.000000	0.000000
1	0.564760	0.564190	0.001011
2	0.399140	0.398942	0.000496
3	0.325840	0.325735	0.000322
4	0.282170	0.282095	0.000267
5	0.252360	0.252313	0.000185
6	0.230370	0.230329	0.000176
7	0.213270	0.213244	0.000124
8	0.199500	0.199471	0.000145
9	0.188080	0.188063	0.000089
10	0.178430	0.178412	0.000099
11	0.170130	0.170110	0.000120
12	0.162880	0.162868	0.000077
13	0.156490	0.156478	0.000076
14	0.150800	0.150786	0.000093
15	0.145680	0.145673	0.000047
16	0.141060	0.141047	0.000089
17	0.136840	0.136836	0.000029
18	0.132990	0.132981	0.000069
19	0.129440	0.129434	0.000047
20	0.126160	0.126157	0.000027
21	0.123120	0.123116	0.000030
22	0.120290	0.120286	0.000036
23	0.117650	0.117642	0.000071
24	0.115170	0.115165	0.000046
25	0.112840	0.112838	0.000018
26	0.110650	0.110647	0.000030
27	0.108580	0.108578	0.000015
28	0.106630	0.106622	0.000077
29	0.104770	0.104767	0.000025
30	0.103010	0.103006	0.000034

2.3 Problem 1c: Diffusion in a Partially Preloaded Semi-Infinite Slab (Val-1c)

This problem models a semi-infinite slab with the first 10 meters preloaded to a uniform concentration. The concentration at the free surface is set to zero for time, $t \geq 0$ sec, when the pre-loaded inventory is allowed to diffuse out the surface and through the slab. No traps are assumed to be present. Comparisons will be made between TMAP7 and analytical values for

concentration histories at two locations: one in the initially unloaded region of the slab, at $x = 12$ m, and one near the surface, $x = 0.5$ m. A third is made at the end of the preloaded region.

By analogy with Carslaw and Jaeger⁷ the concentration as a function of space and time is

$$C = \frac{C_o}{2} \left[2\operatorname{erf}\left(\frac{x}{2\sqrt{Dt}}\right) - \operatorname{erf}\left(\frac{x-h}{2\sqrt{Dt}}\right) - \operatorname{erf}\left(\frac{h+x}{2\sqrt{Dt}}\right) \right] \quad (11)$$

where

h = thickness of pre-loaded region in the slab (10 m)

C_o = concentration of pre-loaded section (1.0 atoms/m³)

D = diffusion coefficient (1.0 m²/sec)

Results for the concentration history at $x = 12$ m can be seen in Table 5. The variance for this problem only exceeded 2% near the peak. Elsewhere it was much less. That could be improved by judicious choice of problem parameters.

Table 5. Concentration history at $x = 12$ meters.

Time (s)	TMAP7	Theory	Variance
0	0.00000	0.00000	0.00000
5	0.26112	0.26340	-0.00865
10	0.32378	0.32007	0.01159
15	0.33563	0.32906	0.01995
20	0.32524	0.31839	0.02152
25	0.30584	0.29989	0.01983
30	0.28373	0.27905	0.01679
35	0.26176	0.25832	0.01333
40	0.24116	0.23877	0.01000
45	0.22234	0.22081	0.00694
50	0.20536	0.20450	0.00418
55	0.19014	0.18979	0.00183
60	0.17650	0.17655	-0.00028
65	0.16428	0.16463	-0.00211
70	0.15331	0.15388	-0.00371
75	0.14345	0.14418	-0.00505
80	0.13455	0.13540	-0.00626
85	0.12650	0.12743	-0.00729
90	0.11920	0.12018	-0.00816
95	0.11256	0.11357	-0.00888
100	0.10650	0.10752	-0.00951

The next comparison for this model is at $x = 0.5$ m, the closest node to the surface. The variance for this problem was less than 2 % for all times except when the value approached zero. Reducing the time step and node spacing could decrease the variance at the expense of calculation time required. These values can be seen in Table 6.

Table 6. Concentration at $x = 0.5$ meters

Time (s)	TMAP7	Theory	Variance
0	0.00000	0.00000	0.00000
5	0.12687	0.12475	0.01699
10	0.08249	0.08164	0.01038
15	0.05953	0.05893	0.01005
20	0.04539	0.04491	0.01062
25	0.03603	0.03560	0.01199
30	0.02947	0.02907	0.01373
35	0.02468	0.02431	0.01527
40	0.02104	0.02070	0.01626
45	0.01820	0.01790	0.01648
50	0.01593	0.01568	0.01597
55	0.01409	0.01388	0.01490
60	0.01256	0.01240	0.01334
65	0.01129	0.01116	0.01144
70	0.01021	0.01012	0.00938
75	0.00929	0.00923	0.00730
80	0.00850	0.00846	0.00524
85	0.00782	0.00779	0.00324
90	0.00722	0.00721	0.00136
95	0.00669	0.00669	-0.00038
100	0.06224	0.00624	8.98023

The last comparison is made at $x = h$. For this case, Equation (11) reduces to

$$C = \frac{C_o}{2} \left[2\operatorname{erf}\left(\frac{h}{2\sqrt{Dt}}\right) - \operatorname{erf}\left(\frac{h+x}{2\sqrt{Dt}}\right) \right]. \quad (12)$$

The variance between the values obtained from TMAP7 and Equation (12), has the largest values at times, $t \leq 20$ sec. For all other times, the variance is less than 0.1 %. Again, the variance may be reduced with judicious selection of problem definition parameters. The comparison of TMAP7 calculated values with theory may be seen in Table 7.

Table 7. Concentration at $x = 10$ meters

Time (s)	TMAP7	Theory	Variance
0	0.50000	0.50000	0.00000
5	0.49862	0.49843	0.00037
10	0.47585	0.47465	0.00253
15	0.43574	0.43224	0.00810
20	0.39160	0.38694	0.01204
25	0.34979	0.34504	0.01378
30	0.31246	0.30821	0.01379
35	0.27993	0.27642	0.01270
40	0.25185	0.24912	0.01097
45	0.22768	0.22567	0.00891
50	0.20683	0.20544	0.00675

Time (s)	TMAP7	Theory	Variance
55	0.18877	0.18791	0.00458
60	0.17306	0.17263	0.00249
65	0.15933	0.15925	0.00052
70	0.14727	0.14746	-0.00126
75	0.13662	0.13702	-0.00289
80	0.12716	0.12773	-0.00442
85	0.11874	0.11942	-0.00573
90	0.11119	0.11196	-0.00687
95	0.10441	0.10524	-0.00789
100	0.09828	0.09915	-0.00879

2.4 Problem 1d: Permeation Problem with Trapping (Val-1da, Val-1db, Val-1dc)

The following three models simulate diffusion through a slab in which traps are operational. The three trapping regimes demonstrated are an effective diffusivity trap, a strong trap, and a set of three traps in the effective diffusivity range with different trap strengths. The diffusion boundary conditions for this set of problems are fixed-concentration or *sconc*, with one surface kept at a constant non-zero concentration and the other set at zero concentration. Initially, the slab is empty. Validation criteria for these problems will be the comparison of the flux and breakthrough times for each of the models with idealizations. The breakthrough time of the flux may have one of two limiting values, which depend on whether the trapping is in the effective diffusivity or strong-trapping regime. A trapping parameter⁸ is defined by

$$\varsigma = \frac{\lambda^2 \nu}{D_o \rho} \exp\left(\frac{E_d - \varepsilon}{kt}\right) + \frac{c}{\rho} \quad (13)$$

where

- λ = lattice parameter (assume 3.162×10^{-8} m)
- ν = Debye frequency (1×10^{13} s⁻¹)
- ρ = trapping site fraction (0.1)
- D_o = diffusivity pre-exponential (1 m²/sec)
- E_d = diffusion activation energy
- ε = trap energy
- k = Boltzmann's constant
- T = temperature (1000 K)
- c = surface dissolved gas atom fraction. (0.0001)

The determining value for which regime is dominant is the relation of ς to c/ρ . If $\varsigma \gg c/\rho$, then the effective diffusivity regime applies, and the flux transient is identical to the standard diffusion transient, but with the diffusivity replaced by an effective diffusivity,

$$D_{eff} = \frac{D}{1 + \sum_i \frac{1}{\zeta_i}} \quad (14)$$

In this limit, the breakthrough time, defined as the intersection of the steepest tangent of the diffusion transient with the time axis, will be

$$\tau_{b_e} = \frac{l^2}{2\pi^2 D_{eff}} \quad (15)$$

where

l = thickness of slab (1 m)

D = diffusivity of gas (1 m²/s).

The permeation transient is then given by

$$J_p = \frac{c_o D}{l} \left[1 + 2 \sum_{m=1}^{\infty} (-1)^m \exp \left(-m^2 \frac{t}{2\tau_{b_e}} \right) \right] \quad (16)$$

where τ_{b_e} is as defined in Equation (15).

2.4.1 Effective Diffusivity Trap (Val-1da)

The first example is the case where a single trap is in the effective diffusivity limit. The ratio ε/k (see Equation (13)) was taken as 100, to give a value of $\zeta = 101$ c/ρ . TMAP7's breakthrough time was found numerically by using a three-point differentiation method given by Fogler⁹ to find the steepest slope.

$$\left(\frac{dC_A}{dt} \right)_{t_i} = \frac{1}{2\Delta t} [C_{A(i+1)} - C_{A(i-1)}] \approx m \quad (17)$$

Then, the point where the slope was the steepest was used with the slope at that point to find the intersection with the time axis. This was computed to be 0.5999 seconds. The analytical breakthrough time was calculated to be 0.611 seconds. The variance between theoretical values of the permeation flux and those calculated by TMAP7 using this model is less than 2%, for most times, as shown in Figure 4. Variance would be less for, say, $\zeta = 1000$ c/ρ . The permeation curve where no trapping is present is also shown in Figure 4 to illustrate the retarding of the permeation curve by a trap.

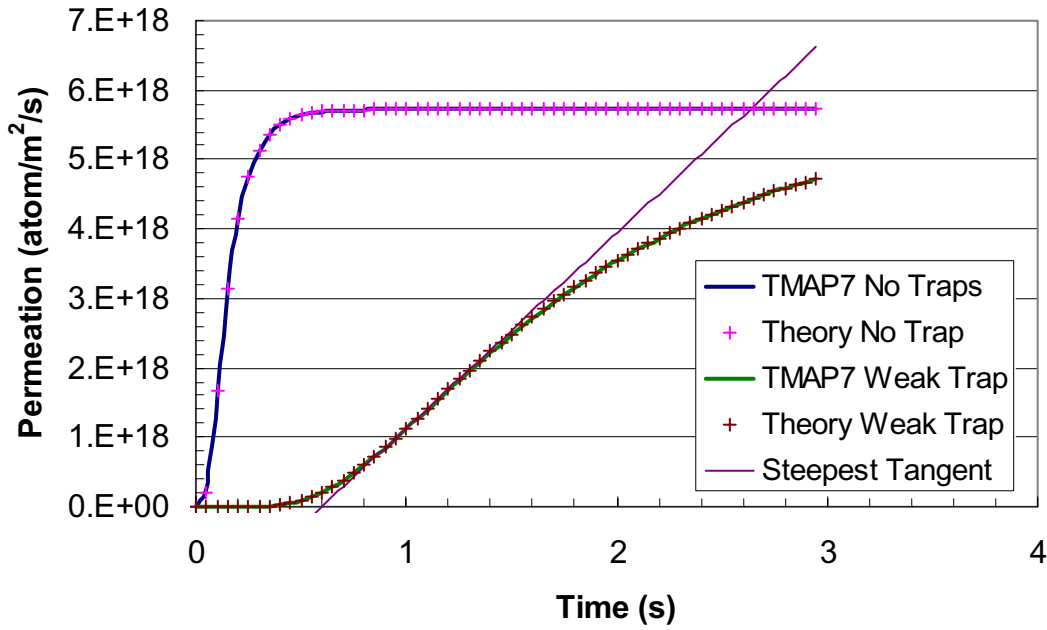


Figure 4. Effective-diffusivity, single trap (Val-1da, Val-1da1).

2.4.2 Strong Trap (Val-1db)

In the second model, $\zeta \ll c/\rho$, is applied to obtain a strong trapping regime. In this regime, no permeation occurs until essentially all the traps have been filled. Then the permeation rapidly turns on to its steady state value. This is due to the relatively low release of trapped atoms. The breakthrough time is given by

$$\tau_{b_d} = \frac{l^2 \rho}{2c_o D} \quad (18)$$

where c_o , ρ , l , and D are defined as in the first model. The value of ε/k is taken to be 15,000 K, to give $\zeta = c/\rho$. The only difference in the input file between the first and second models is this parameter and a larger time step. The breakthrough time in the strong trapping regime was taken as the first time that the permeation was at its steady state value. This occurred at 511 seconds. The estimated breakthrough time from Eq. (18) is 500 seconds. The permeation curve for this model can be seen in Figure 5.

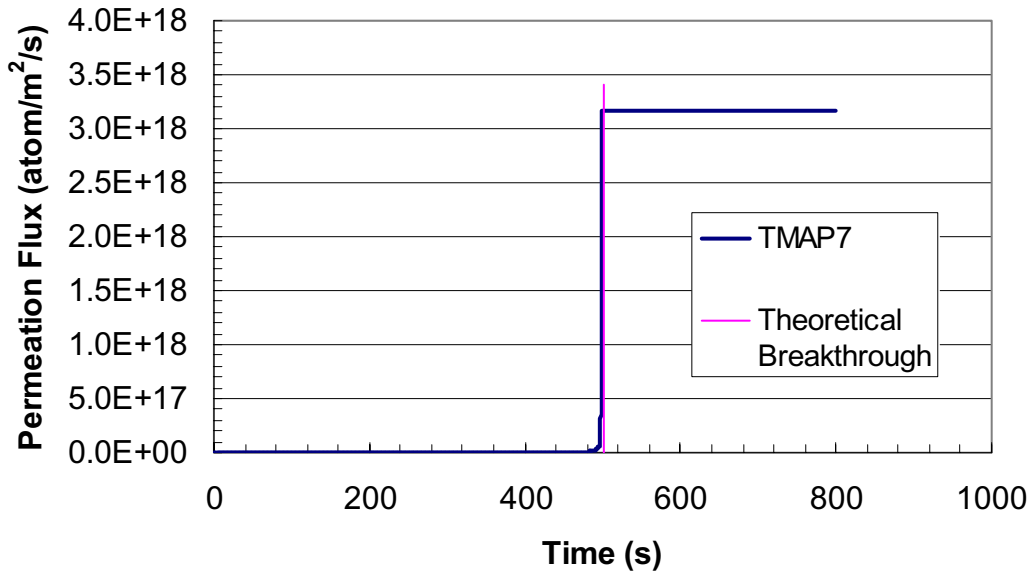


Figure 5. Permeation for strong-trapping regime (Val-1db)

2.4.3 Multiple Trap (Val-1dc)

The last problem modeled in this section demonstrates the effects of multiple traps. This feature is new to TMAP7. To illustrate TMAP7's capabilities to allow for multiple traps, three traps that are relatively weak are assumed to be active in a slab. The parameters of the first trap are the same as the trap in the effective diffusivity limit, first model. The second and third traps vary by having trap concentrations of 0.15 and 0.20 atom fractions and the values of ε/k chosen to be 500 K and 800 K, respectively. These values give the following values for ζ :

Trap 1: 181.97 c/ρ

Trap 2: 122.31 c/ρ

Trap 3: 90.87 c/ρ .

The effective diffusivity was calculated from Equation (14), $D_{eff} = 0.01242 \text{ m}^2/\text{sec}$, and the breakthrough time was calculated from Equation (15) to be 4.08 sec. TMAP7's calculated breakthrough time was 4.00 sec. The permeation curves that were calculated using Equation (16) are compared with TMAP7 results in Figure 6. The graphs for the theoretical flux and the calculated flux are nearly identical.

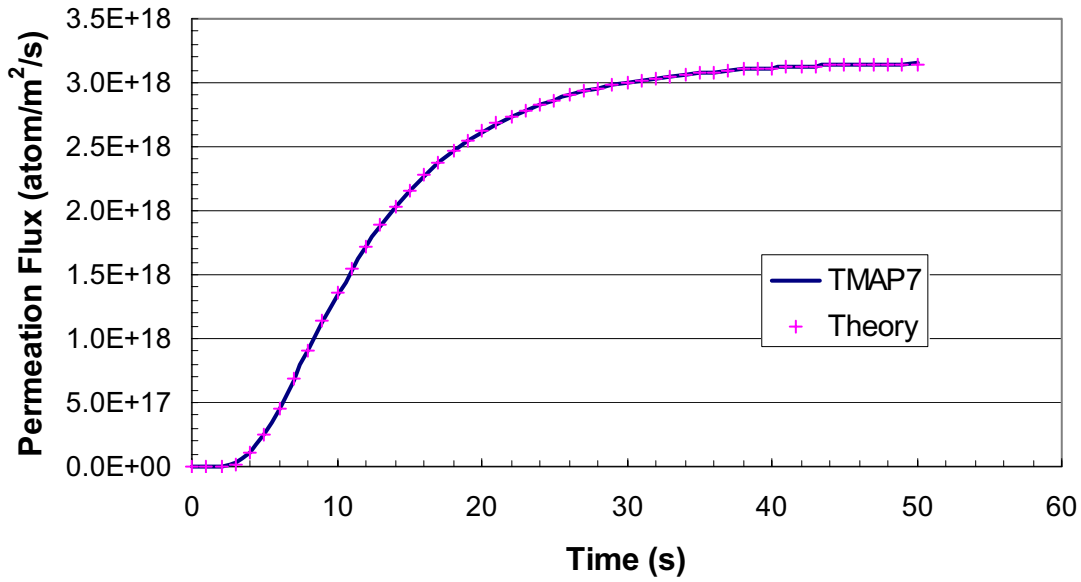


Figure 6. Permeation curve for slab with multiple traps (Val-1dc).

2.5 Problem 1e: Diffusion with Composite Material Layers (Val-1e)

A composite structure of PyC and SiC is modeled with a constant concentration boundary condition on the free surface of the PyC and a zero-concentration boundary on the free surface of the SiC. The concentration profile in steady state is to be analyzed. The steady-state solution for the PyC is given in Eq. (19),

$$C = C_o \left[1 + \frac{x}{a} \left(\frac{D_{PyC} l}{l D_{PyC} + a D_{SiC}} - 1 \right) \right] \quad (19)$$

while the concentration profile for the SiC is given by

$$C = C_o \left(\frac{a + l - x}{l} \right) \left(\frac{D_{PyC} l}{l D_{PyC} + a D_{SiC}} \right) \quad (20)$$

where

a = thickness of the PyC layer (33 μm)

l = thickness of the SiC layer (66 μm)

C_o = the concentration at the surface (3.0537×10^{25} atoms/m³)

S_a = Solubility of both species was taken as 1.0 (units arbitrary)

The values for the diffusivity were taken as constants, $D_{PyC} = 1.274 \times 10^{-7}$ m²/sec and $D_{SiC} = 2.622 \times 10^{-11}$ m²/sec. The variance for this problem does not exceed 0.004%. The comparison of Eqs. (19) and (20) with TMAP7's values can be seen in Table 8.

Table 8. Steady-State Concentration Profile in Composite Slab

Depth (m)	TMAP7	Theory	Variance
0.00E+00	3.0537E+25	3.0537E+25	0.00E+00
1.50E-06	3.0537E+25	3.0537E+25	-4.68E-06
5.50E-06	3.0536E+25	3.0536E+25	1.56E-05
1.05E-05	3.0536E+25	3.0536E+25	8.27E-09
1.55E-05	3.0536E+25	3.0536E+25	-1.56E-05
2.05E-05	3.0535E+25	3.0535E+25	1.58E-06
2.55E-05	3.0535E+25	3.0535E+25	-1.40E-05
3.05E-05	3.0534E+25	3.0534E+25	3.14E-06
3.30E-05	3.0534E+25	3.0534E+25	-4.65E-06
3.30E-05	3.0534E+25	3.0534E+25	-4.65E-06
3.83E-05	2.8105E+25	2.8105E+25	1.01E-06
4.88E-05	2.3247E+25	2.3247E+25	1.59E-05
5.93E-05	1.8390E+25	1.8390E+25	-1.58E-05
6.98E-05	1.3532E+25	1.3532E+25	3.74E-06
8.03E-05	8.6744E+24	8.6744E+24	-9.85E-07
9.08E-05	3.8167E+24	3.8167E+24	8.45E-06
9.75E-05	6.9395E+23	6.9395E+23	1.90E-06
9.90E-05	0.0000E+00	0.0000E+00	0.00E+00

Demonstration of transient agreement with theory may also be shown by examining the concentration history at a point 7.5 μm into the SiC layer as a function of time given that, initially, both PyC and SiC were empty of gas. The transient solution for concentration in the SiC side of the composite slab is

$$C = C_o \left\{ \frac{D_{PyC}(l-x)}{lD_{PyC} + aD_{SiC}} - 2 \sum_{n=1}^{\infty} \frac{\sin(a\lambda_n)\sin(kl\lambda_n)\sin[k(l-x)\lambda_n]}{\lambda_n [a \sin(kl\lambda_n) + l \sin^2(a\lambda_n)]} \exp(-D_{PyC}\lambda_n^2 t) \right\} \quad (21)$$

where

a = thickness of PyC (33 μm)

l = Thickness of SiC (66 μm)

$$k = \sqrt{\frac{D_{PyC}}{D_{SiC}}} = 69.7036$$

and the λ_n are the roots of

$$\tan(\lambda a) + k \tan(k \lambda l) = 0 \quad (22)$$

Figure 7 shows the graphical comparison, and Table 9 lists discrete values and variance. The series in Eq. (21) was evaluated with 19 terms such that the contribution of the last term was less than 1.0E-11 for all times. The fit improves with finer spatial mesh.

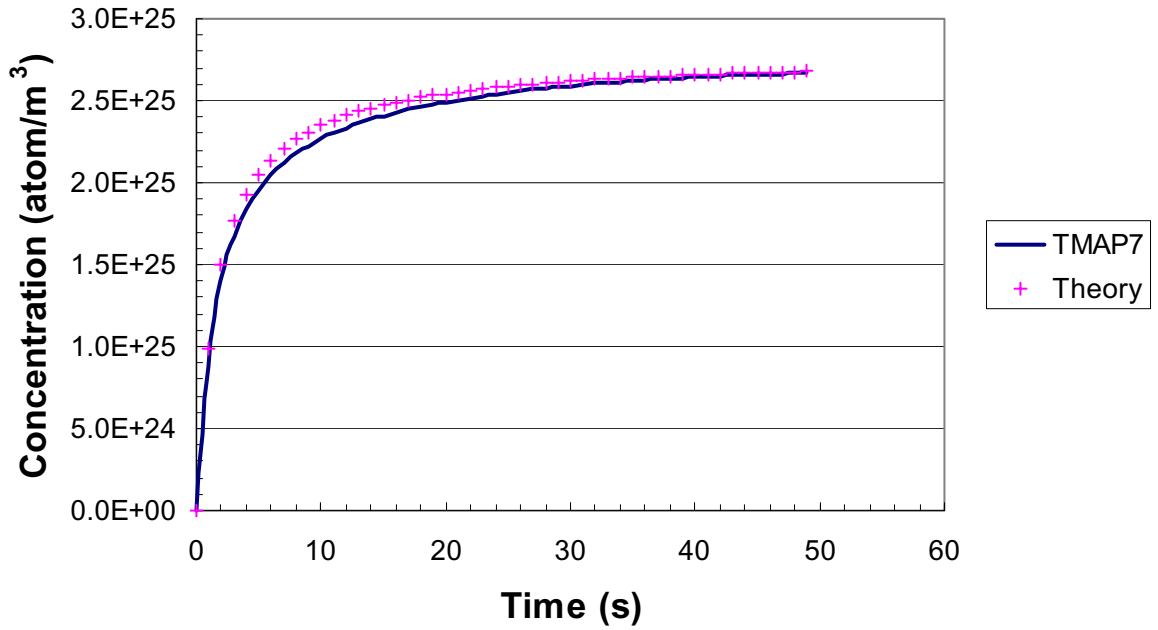


Figure 7. Comparison of TMAP7 with theoretical solution for transient concentration at a location 7.5 μm deep in the SiC layer of a composite slab of PyC and SiC (Val-1e).

Table 9. Variance for transient solution in composite slab.

Time (s)	TMAP7	Theory	Variance
0	0.00E+00	0.00E+00	0.000000
1	8.81E+24	9.92E+24	0.111677
2	1.40E+25	1.50E+25	0.067949
3	1.67E+25	1.76E+25	0.055549
4	1.84E+25	1.93E+25	0.049649
5	1.96E+25	2.05E+25	0.045956
6	2.05E+25	2.14E+25	0.043165
7	2.12E+25	2.21E+25	0.040838
8	2.18E+25	2.26E+25	0.038742
9	2.23E+25	2.31E+25	0.036767
10	2.27E+25	2.35E+25	0.034865
11	2.30E+25	2.38E+25	0.032995
12	2.33E+25	2.41E+25	0.031260
13	2.36E+25	2.43E+25	0.029555
14	2.39E+25	2.45E+25	0.027897
15	2.41E+25	2.47E+25	0.026365
16	2.43E+25	2.49E+25	0.024870
17	2.45E+25	2.51E+25	0.023453
18	2.46E+25	2.52E+25	0.022090
19	2.48E+25	2.53E+25	0.020817
20	2.49E+25	2.54E+25	0.019579
21	2.51E+25	2.55E+25	0.018465
22	2.52E+25	2.56E+25	0.017358

Time (s)	TMAP7	Theory	Variance
23	2.53E+25	2.57E+25	0.016332
24	2.54E+25	2.58E+25	0.015376
25	2.55E+25	2.59E+25	0.014475
26	2.56E+25	2.59E+25	0.013650
27	2.57E+25	2.60E+25	0.012839
28	2.58E+25	2.61E+25	0.012059
29	2.58E+25	2.61E+25	0.011361
30	2.59E+25	2.62E+25	0.010718
31	2.60E+25	2.62E+25	0.010063
32	2.60E+25	2.63E+25	0.009481
33	2.61E+25	2.63E+25	0.008906
34	2.62E+25	2.64E+25	0.008418
35	2.62E+25	2.64E+25	0.007913
36	2.63E+25	2.65E+25	0.007433
37	2.63E+25	2.65E+25	0.007022
38	2.63E+25	2.65E+25	0.006609
39	2.64E+25	2.66E+25	0.006237
40	2.64E+25	2.66E+25	0.005872

2.6 Problem 1f: Heat Sink/Source Problem

Four heat transfer models were set up to validate the heat transfer capabilities of the TMAP7 code. The four problems solved include (a) heat conduction with generation; (b) conduction modeled as mass transfer, (c) transient conduction and steady state values in a composite structure, and (d) heating of a semi-infinite slab by convection.

2.6.1 Heat conduction with generation (Val-1fa)

To model the first problem, the thermal boundary conditions were set so one surface was adiabatic, while the other was kept at constant temperature. The heat generation in the slab was assumed to be constant throughout. Incropera and DeWitt¹⁰ give the analytical solution for the steady state temperature of this model as

$$T = T_s + \frac{QL^2}{2k} \left(1 - \frac{x^2}{L^2} \right) \quad (23)$$

where

Q = internal heat generation rate (10,000 W/m³)

L = thickness of slab (1.6 m)

k = thermal conductivity (10 W/m K)

T_s = surface temperature (300 K)

A value for thermal mass, the product of material mass density and specific heat, must be added for TMAP7 thermal calculations. In this problem, $\rho c_p = 1$ J/m³K was assumed. Initially, 16 spatial segments were assumed. The variance for this problem was less than 0.2% for distances less than 1.35 m, but it increased as the distance from the adiabatic surface was

increased. To show that this can be reduced with a decrease in the distance between nodes, an additional calculation was performed with 48 spatial segments. The variance was reduced by a factor of approximately 10. The comparison of Eq. (23) with TMAP7 values can be seen in Table 10.

Table 10. Heat Conduction with Generation

Position (m)	Theory	16 Segs	Variance	48 Segs	Variance
0.00	1580.00	1580.00	0.00000	1580.00	0.00000
0.05	1578.75	1580.00	0.00079	1578.90	0.00010
0.15	1568.75	1570.00	0.00080	1568.90	0.00010
0.25	1548.75	1550.00	0.00081	1548.90	0.00010
0.35	1518.75	1520.00	0.00082	1518.90	0.00010
0.45	1478.75	1480.00	0.00085	1478.90	0.00010
0.55	1428.75	1430.00	0.00087	1428.90	0.00010
0.65	1368.75	1370.00	0.00091	1368.90	0.00011
0.75	1298.75	1300.00	0.00096	1298.90	0.00012
0.85	1218.75	1220.00	0.00103	1218.90	0.00012
0.95	1128.75	1130.00	0.00111	1128.90	0.00013
1.05	1028.75	1030.00	0.00122	1028.90	0.00015
1.15	918.75	920.00	0.00136	918.88	0.00014
1.25	798.75	800.00	0.00156	798.88	0.00016
1.35	668.75	670.00	0.00187	668.88	0.00019
1.45	528.75	530.00	0.00236	528.88	0.00025
1.55	378.75	380.00	0.00330	378.89	0.00037
1.60	300.00	300.00	0.00000	300.00	0.00000

2.6.2 Thermal Diffusion Transient (Val-1fb)

The second problem validates the thermal diffusion capability in a slab. The temperature of the left side of the thermal segment was held constant at 400 K while the right side was held at a constant 300 K. The initial temperature in the slab was 300 K. For this example, the thickness, L , was 4 m and the heat production rate was $\dot{Q} = 0$. Mass diffusion was ignored by setting the mobile species concentration to zero and using non-flow boundaries. The analytical solution is given by

$$T(x,t) = T_o + (T_1 - T_o) \left[1 - \frac{x}{L} - \frac{2}{L} \sum_{m=0}^{\infty} \frac{1}{\lambda_m} \sin(\lambda_m x) \exp(-\alpha \lambda_m^2 t) \right] \quad (24)$$

where

$$\lambda_m = m \frac{\pi}{L} \quad (25)$$

and thermal diffusivity is

$$\alpha = \frac{k}{C_p \rho} \quad (26)$$

For the problem analyzed,

$$\alpha = 1.0 \text{ m}^2/\text{s},$$

$$T_o = 300 \text{ K, and}$$

$$T_1 = 400 \text{ K.}$$

The values for Eq. (24) were found using MS Excel. The last term in the summation taken contributed less than 1×10^{-13} of the theoretical value. The agreement between TMAP7 and Eq. (24) is excellent, with the variance less than 1 % for each case tested, and usually much less. The comparison between the values can be seen in Figure 8 for temperature profiles through the slab at 0.1, 0.5, 1.0, and 5.0 seconds.

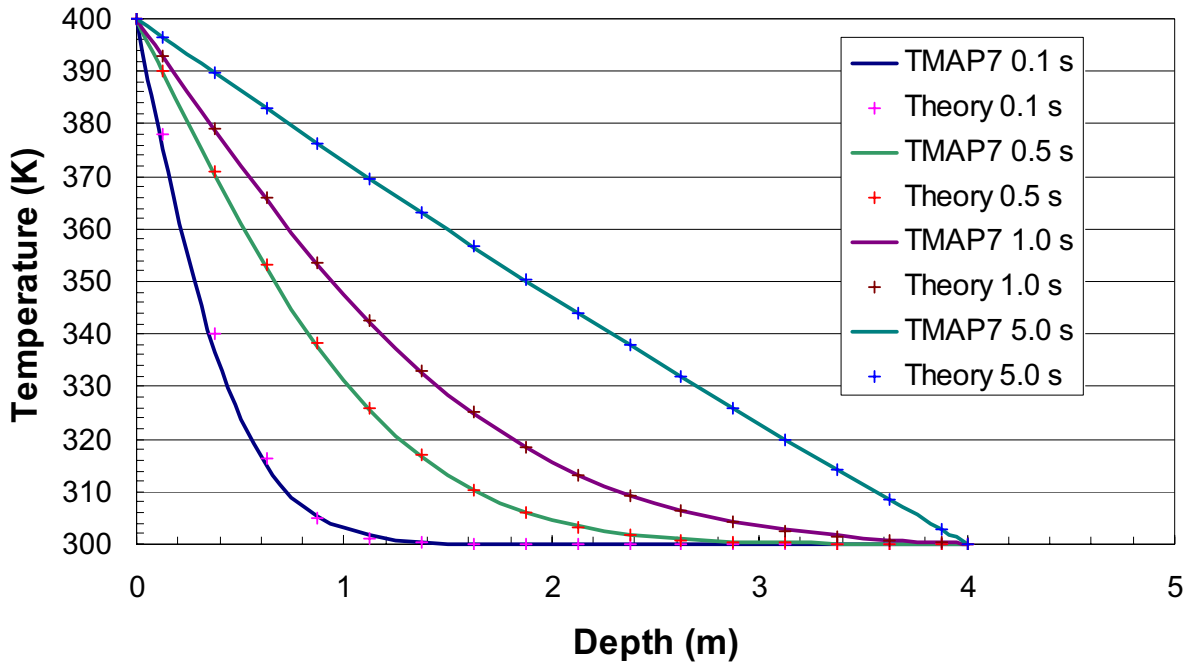


Figure 8. Transient temperature distribution for various times in a slab (Val-1fb).

2.6.3 Conduction in Composite Structure with Constant Surface Temperatures (Val-1fc)

The third heat transfer problem studied was heat transfer through a composite slab with constant surface temperatures. The composite was a 40-cm thick layer of Cu followed by a 40-cm layer of Fe. The temperature of both layers was initially 0 K, but at time $t = 0$, the outside face of the copper was held at 600 K while the outside face of the Fe was maintained at 0 K. This problem was modeled using the heat transfer capability of TMAP7. The computational answers obtained by TMAP7 for both the transient and steady state solutions were compared to values obtained from ABAQUS.¹¹ The ABAQUS code was setup and run by R. G. Ambrosek. ABAQUS is a heat transfer program that has been validated for both transient and steady state solutions. The transient solution was compared at a constant time and constant distance. The constant time comparison between ABAQUS and TMAP7 was made at time, $t = 150$ sec. The

variance in this comparison grows with increasing distance. This may be due to the time interval on both programs being larger than needed, or round-off error from the printed values. These values can be seen in Table 11.

Table 11. Temperature distribution in composite structure at $t = 150$ seconds.

Depth (m)	ABAQUS	TMAP7	Variance
0.00	600.000	600.000	0.00000
0.01	574.400	574.370	-0.00005
0.03	523.600	523.400	-0.00038
0.05	473.600	473.300	-0.00063
0.07	425.100	424.630	-0.00111
0.09	378.400	377.880	-0.00137
0.11	334.100	333.500	-0.00180
0.13	292.500	291.850	-0.00222
0.15	253.900	253.200	-0.00276
0.17	218.500	217.770	-0.00334
0.19	186.400	185.670	-0.00392
0.21	157.700	156.930	-0.00488
0.23	132.200	131.510	-0.00522
0.25	110.000	109.320	-0.00618
0.27	90.790	90.199	-0.00651
0.29	74.480	73.975	-0.00678
0.31	60.860	60.439	-0.00692
0.33	49.690	49.374	-0.00636
0.35	40.770	40.564	-0.00505

The values were also compared at $x = 0.09$ m, at 5 second intervals from time $t = 0$ to 150 sec. The variance is initially large, but reduces as the time increases. The initially large variance may be due to the same factors of spatial resolution and time step size mentioned earlier. These results can be seen in Table 12.

The steady-state solution for this problem was compared to the analytical solution in addition to the ABAQUS answer. To solve for the steady state solution for this problem, the heat flux is given by

$$q'' = \frac{T_{S_A} - T_{S_B}}{\frac{L_A}{k_A} + \frac{L_B}{k_B}} \quad (27)$$

where

T_{Si} = Temperature of surface i, left (A) and right (B),

L_i = Length of segment i

k_i = thermal conductivity of segment i.

Table 12. Temperature history in composite structure at $x = 0.09$ meters

Time (s)	TMAP7	ABAQUS	Variance
0	0.00	0.00	0.00000
5	6.50	10.11	0.35692
10	37.03	43.47	0.14817
15	75.11	81.42	0.07745
20	110.42	115.90	0.04728
25	140.99	145.60	0.03166
30	167.17	171.10	0.02297
35	189.68	193.10	0.01771
40	209.20	212.10	0.01367
45	226.29	228.90	0.01140
50	241.39	243.70	0.00948
55	254.84	256.90	0.00802
60	266.91	268.70	0.00666
65	277.82	279.50	0.00601
70	287.74	289.36	0.00560
75	296.81	298.30	0.00499
80	305.14	306.40	0.00411
85	312.83	314.00	0.00373
90	319.95	321.00	0.00327
95	326.57	327.60	0.00314
100	332.74	333.70	0.00288
105	338.52	339.40	0.00259
110	343.94	344.80	0.00249
115	349.04	349.80	0.00217
120	353.85	354.60	0.00212
125	358.39	359.10	0.00198
130	362.70	363.40	0.00193
135	366.78	367.40	0.00169
140	370.66	371.30	0.00172
145	374.36	374.90	0.00144
150	377.88	378.40	0.00137

For the solution to be at steady state, the flux in and out of any section of the slab must be equal. The temperature at the interface can be found by setting the flux through A equal to the flux through B .

$$\frac{T_{S_A} - T_I}{\frac{L_A}{k_A}} = \frac{T_I - T_{S_B}}{\frac{L_B}{k_B}} \quad (28)$$

where

T_I = temperature of interface,

$k_A = 401 \text{ W /m K}$,

$$k_B = 80.2 \text{ W /m K},$$

$$L_A = L_B = 0.4 \text{ meters},$$

$$T_{S_A} = 600 \text{ K}$$

$$T_{S_B} = 0 \text{ K}$$

From Eq. (28), the interface temperature is found to be $T_I = 500 \text{ K}$. The temperature profile for conduction in a steady state, constant physical properties, is linear. TMAP7 was run to a time of 150,000 seconds. Steady state was assured by there being no difference at all in the last two temperature profiles. The theoretical temperature profile of A and B can be found through linear interpolation. The steady-state temperatures predicted by TMAP7, ABAQUS, and the analytical solution were found to be identical. These values can be seen in Table 13.

Table 13. Steady-state temperature (K) distribution for composite structure

Depth (m)	TMAP7	ABAQUS	Theory
0.00	600.0	600.0	600.0
0.01	597.5	597.5	597.5
0.03	592.5	592.5	592.5
0.05	587.5	587.5	587.5
0.07	582.5	582.5	582.5
0.09	577.5	577.5	577.5
0.11	572.5	572.5	572.5
0.13	567.5	567.5	567.5
0.15	562.5	562.5	562.5
0.17	557.5	557.5	557.5
0.19	552.5	552.5	552.5
0.21	547.5	547.5	547.5
0.23	542.5	542.5	542.5
0.25	537.5	537.5	537.5
0.27	532.5	532.5	532.5
0.29	527.5	527.5	527.5
0.31	522.5	522.5	522.5
0.33	517.5	517.5	517.5
0.35	512.5	512.5	512.5
0.37	507.5	507.5	507.5
0.39	502.5	502.5	502.5
0.40	500.0	500.0	500.0
0.41	487.5	487.5	487.5
0.43	462.5	462.5	462.5
0.45	437.5	437.5	437.5
0.47	412.5	412.5	412.5
0.49	387.5	387.5	387.5
0.51	362.5	362.5	362.5
0.53	337.5	337.5	337.5
0.55	312.5	312.5	312.5
0.57	287.5	287.5	287.5
0.59	262.5	262.5	262.5
0.61	237.5	237.5	237.5

Depth (m)	TMAP7	ABAQUS	Theory
0.63	212.5	212.5	212.5
0.65	187.5	187.5	187.5
0.67	162.5	162.5	162.5
0.69	137.5	137.5	137.5
0.71	112.5	112.5	112.5
0.73	87.5	87.5	87.5
0.75	62.5	62.5	62.5
0.77	37.5	37.5	37.5
0.79	12.5	12.5	12.5
0.80	0.0	0.0	0.0

2.6.4 Convective Heating (Val-1fd)

The fourth heat transfer problem modeled was the heating of a semi-infinite slab by convection at the boundary. The slab was initially configured with a constant temperature of 100 K throughout the slab. A convection boundary was then activated at the surface for time, $t \geq 0$ sec. Incorpera and DeWitt¹⁰ give for the solution

$$T(x,t) = T_i + (T_\infty - T_i) \left\{ \left[\operatorname{erfc} \left(\frac{x}{2\sqrt{t\alpha}} \right) \right] - \left[\exp \left(\frac{hx}{k} + \frac{h^2 t \alpha}{k^2} \right) \right] \left[\operatorname{erfc} \left(\frac{x}{2\sqrt{t\alpha}} + \frac{h\sqrt{t\alpha}}{k} \right) \right] \right\} \quad (29)$$

where

T_i = initial temperature (100 K)

T_∞ = temperature of enclosure (500 K)

h = conduction coefficient (200 W/m² K)

k = thermal conductivity (401 W/m K)

α = thermal diffusivity (1.17 x 10⁻⁴ m²/s)

The depth x of 5 cm was used for comparison. Values of the complimentary error function were computed using a series expansion in MS ExcelTM. The last term computed contributed less than 1.0×10^{-20} . The variance between Eq. (29) and TMAP7 was less than 0.2%, for all times greater than 30 sec, as can be seen in Table 14. A graphical comparison can be seen in Figure 9.

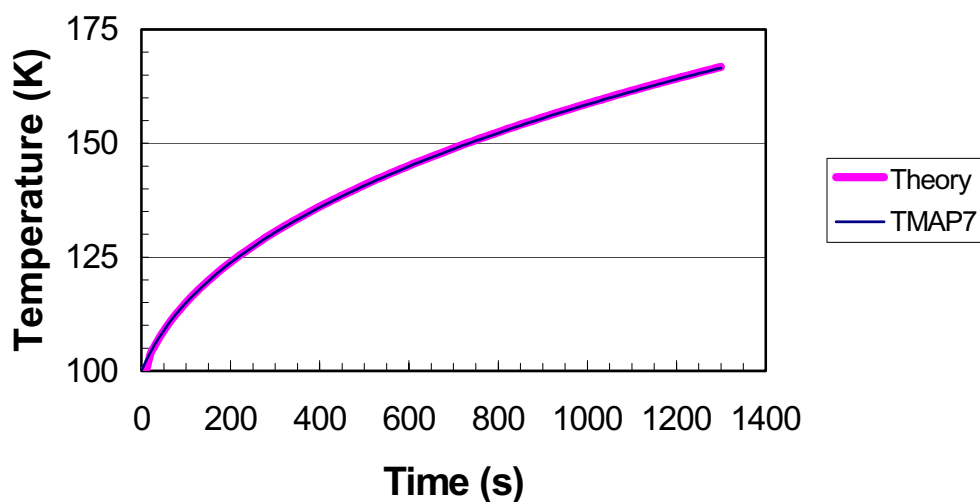


Figure 9. Convective heating at depth 5 cm in a semi-infinite slab (Val-1fd).

Table 14. Heating of Semi-Infinite Slab by Convection

Time (s)	TMAP7	Theory	Variance
0	100.000	100.000	0.000000
10	102.140	100.000	0.021400
20	104.060	103.639	0.004063
30	105.800	105.549	0.002376
40	107.390	107.252	0.001286
50	108.860	108.795	0.000600
60	110.230	110.211	0.000169
70	111.510	111.527	-0.000148
80	112.720	112.758	-0.000333
90	113.860	113.918	-0.000508
100	114.940	115.017	-0.000673
110	115.980	116.064	-0.000724
120	116.970	117.064	-0.000806
130	117.920	118.023	-0.000876
140	118.840	118.945	-0.000887
150	119.720	119.834	-0.000953
160	120.580	120.693	-0.000934
170	121.410	121.524	-0.000934
180	122.210	122.329	-0.000973

2.7 Problem 1g: Enclosure Reaction Problems

Three problems were solved in TMAP7 to test its capability to handle enclosure reactions. The first model is a simple forward reaction with two reactants forming one product. In the first model, the reactants start in their stoichiometric ratio. The second problem varies from the first in that the concentrations of the reactants vary from their stoichiometric ratio. The third problem examines a series reaction.

2.7.1 Simple Forward Reactions (Val-1ga and Val-1gb)

The first and second problems consider the simple chemical reaction



The rate at which the concentrations change (rate of reaction) is assumed first order with respect to the concentrations of A and B. The rate coefficient, K_r , is a constant for the reaction and has no spatial or time dependence. The simple forward reaction rate

$$\frac{dC_{AB}}{dt} = R_c = K_r C_A C_B \quad (31)$$

is positive if AB is produced and negative if AB is consumed in the reaction. This may also be written

$$\left(\frac{dC_{AB}}{C_{AB} - C_{A_0}} \right) \left(\frac{dC_{AB}}{C_{AB} - C_{B_0}} \right) = K_r dt \quad (32)$$

The solution for this problem is¹²

$$C_{AB} = C_{B_0} \frac{1 - \exp[K_R t (C_{B_0} - C_{A_0})]}{1 - \frac{C_{B_0}}{C_{A_0}} \exp[K_R t (C_{B_0} - C_{A_0})]} \quad (33)$$

where

C_{AB} = concentration of species AB

C_{A_0} = initial concentration of species A

C_{B_0} = initial concentration of species B

If $C_{A_0} = C_{B_0}$, Eq. (33) can be simplified to

$$C_{AB} = C_{A_0} - \frac{1}{\frac{1}{C_{A_0}} + K_R t}. \quad (34)$$

The analytical solutions of Eqs. (33) and (34) were found and compared to the values obtained from TMAP7. Eq. (34) was solved and compared to TMAP7 for problem Val-1ga, and Eq. (33) was compared to TMAP7 for problem Val-1gb. These results are listed in Tables 15 and 16, respectively. Figure 10 shows a graphical comparison of the two cases. The variance in each of the two cases drops below 0.2% for time, $t \geq 2$ sec.

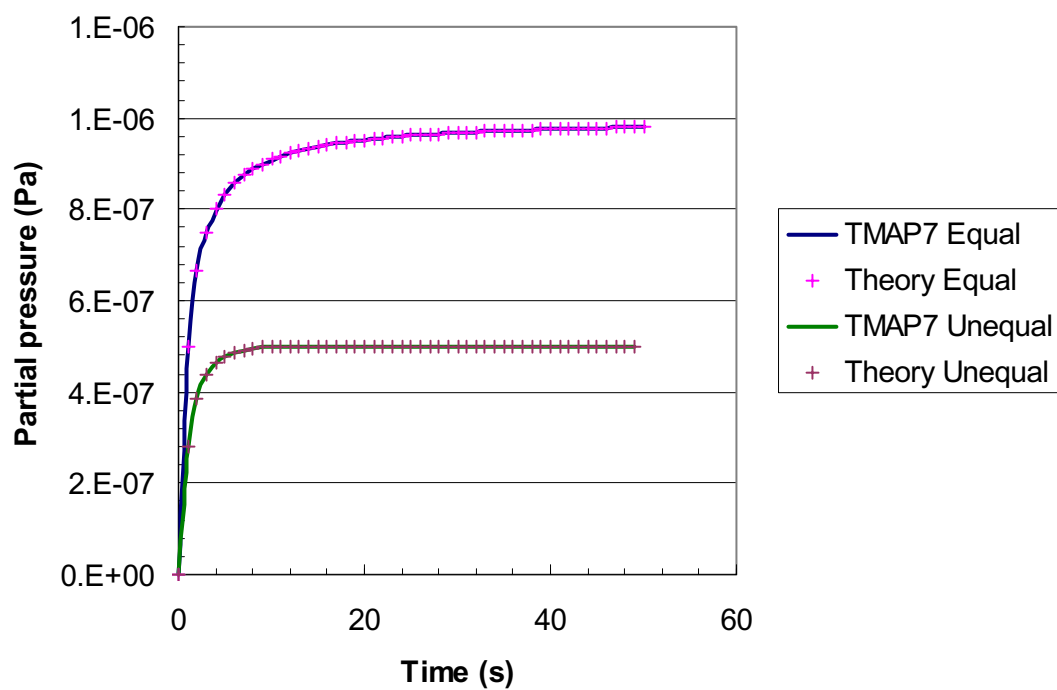


Figure 10. Production of AB from A and B under assumptions of equal and unequal initial reactant concentrations (Val-1ga/Val-1gb).

Table 15. Partial pressure (Pa) of product for equal starting concentrations.

Time (s)	TMAP7	Theory	Variance
0	0.00E+00	0.00E+00	0.0000
1	4.98E-07	5.00E-07	-0.0037
2	6.65E-07	6.67E-07	-0.0020
3	7.49E-07	7.50E-07	-0.0013
4	7.99E-07	8.00E-07	-0.0010
5	8.33E-07	8.33E-07	-0.0008
6	8.57E-07	8.57E-07	-0.0006
7	8.75E-07	8.75E-07	-0.0005
8	8.88E-07	8.89E-07	-0.0005
9	9.00E-07	9.00E-07	-0.0004
10	9.09E-07	9.09E-07	-0.0004
11	9.16E-07	9.17E-07	-0.0003
12	9.23E-07	9.23E-07	-0.0003
13	9.28E-07	9.29E-07	-0.0003
14	9.33E-07	9.33E-07	-0.0003
15	9.37E-07	9.37E-07	-0.0003
16	9.41E-07	9.41E-07	-0.0002
17	9.44E-07	9.44E-07	-0.0002
18	9.47E-07	9.47E-07	-0.0002
19	9.50E-07	9.50E-07	-0.0002

Time (s)	TMAP7	Theory	Variance
20	9.52E-07	9.52E-07	-0.0002
21	9.54E-07	9.55E-07	-0.0002
22	9.56E-07	9.57E-07	-0.0002
23	9.58E-07	9.58E-07	-0.0002
24	9.60E-07	9.60E-07	-0.0002
25	9.61E-07	9.62E-07	-0.0002
26	9.63E-07	9.63E-07	-0.0002
27	9.64E-07	9.64E-07	-0.0002
28	9.65E-07	9.66E-07	-0.0002
29	9.66E-07	9.67E-07	-0.0002
30	9.68E-07	9.68E-07	-0.0002
31	9.69E-07	9.69E-07	-0.0002
32	9.70E-07	9.70E-07	-0.0002
33	9.70E-07	9.71E-07	-0.0002
34	9.71E-07	9.71E-07	-0.0002
35	9.72E-07	9.72E-07	-0.0002
36	9.73E-07	9.73E-07	-0.0002
37	9.74E-07	9.74E-07	-0.0002
38	9.74E-07	9.74E-07	-0.0002
39	9.75E-07	9.75E-07	-0.0002
40	9.75E-07	9.76E-07	-0.0002
41	9.76E-07	9.76E-07	-0.0002
42	9.77E-07	9.77E-07	-0.0002
43	9.77E-07	9.77E-07	-0.0002
44	9.78E-07	9.78E-07	-0.0002
45	9.78E-07	9.78E-07	-0.0002
46	9.79E-07	9.79E-07	-0.0002
47	9.79E-07	9.79E-07	-0.0002
48	9.79E-07	9.80E-07	-0.0002
49	9.80E-07	9.80E-07	-0.0002
50	9.80E-07	9.80E-07	-0.0002

Table 16. Partial pressure (Pa) of product for reaction with unequal starting concentrations.

Time (s)	TMAP7	Theory	Variance
0	0.00E+00	0.00E+00	0.0000
1	2.82E-07	2.82E-07	-0.0005
2	3.87E-07	3.87E-07	-0.0003
3	4.37E-07	4.37E-07	-0.0003
4	4.64E-07	4.64E-07	-0.0002
5	4.79E-07	4.79E-07	-0.0002
6	4.87E-07	4.87E-07	-0.0002
7	4.92E-07	4.92E-07	-0.0002
8	4.95E-07	4.95E-07	-0.0002
9	4.97E-07	4.97E-07	-0.0001
10	4.98E-07	4.98E-07	-0.0002
11	4.99E-07	4.99E-07	-0.0002
12	4.99E-07	4.99E-07	-0.0001
13	5.00E-07	5.00E-07	-0.0001
14	5.00E-07	5.00E-07	-0.0001
15	5.00E-07	5.00E-07	-0.0001
16	5.00E-07	5.00E-07	-0.0002
17	5.00E-07	5.00E-07	-0.0001
18	5.00E-07	5.00E-07	-0.0001
19	5.00E-07	5.00E-07	-0.0001
20	5.00E-07	5.00E-07	-0.0001
21	5.00E-07	5.00E-07	-0.0001
22	5.00E-07	5.00E-07	-0.0002
23	5.00E-07	5.00E-07	-0.0001
24	5.00E-07	5.00E-07	-0.0001
25	5.00E-07	5.00E-07	-0.0001
26	5.00E-07	5.00E-07	-0.0001
27	5.00E-07	5.00E-07	-0.0001
28	5.00E-07	5.00E-07	-0.0001
29	5.00E-07	5.00E-07	-0.0001
30	5.00E-07	5.00E-07	-0.0001
31	5.00E-07	5.00E-07	-0.0001
32	5.00E-07	5.00E-07	-0.0001
33	5.00E-07	5.00E-07	-0.0001
34	5.00E-07	5.00E-07	-0.0001
35	5.00E-07	5.00E-07	-0.0001
36	5.00E-07	5.00E-07	-0.0001
37	5.00E-07	5.00E-07	-0.0001
38	5.00E-07	5.00E-07	-0.0001
39	5.00E-07	5.00E-07	-0.0001
40	5.00E-07	5.00E-07	-0.0001

2.7.2 Series Reactions (Val-1gc)

The third problem modeled is a set of reactions in series. The system was configured so that the enclosure initially contained only species A. At time $t \geq 0$, the reactions were allowed to proceed. The reactions that were modeled are



The production rate for each species (negative means consumption) is given by

$$-r'_A = k_1 C_A \quad (36)$$

$$r'_B = k_1 C_A - k_2 C_B \quad (37)$$

$$r'_C = k_2 C_B \quad (38)$$

Fogler¹³ gives the concentrations of A and B as

$$C_A = C_{A_0} \exp(-k_1 t) \quad (39)$$

$$C_B = k_1 C_{A_0} \left(\frac{\exp(-k_1 t) - \exp(-k_2 t)}{k_2 - k_1} \right) \quad (40)$$

where

t = time (sec),

C_{A_0} = initial concentration of A, (2.415×10^{14} atoms/m³).

k_1 = rate constant of reaction 1 (0.0125 s^{-1})

k_2 = rate constant of reaction 2 (0.0025 s^{-1}).

The concentration of C was found by applying a mass balance over the system. From the stoichiometry of this reaction it was found that

$$C_C = C_{A_0} - C_A - C_B . \quad (41)$$

The concentration values of Eqs. (39), (40), and (41) were obtained using MS Excel™ and converted to Pa. These numbers were then compared to the partial pressure values obtained from TMAP7. The variance for the pressures of species A and B are less than 0.03% for all time. The variance of species C, begins at around 0.1%, but continually decreases as the problem time increases. The comparisons for this problem are listed in Table 17. A graphical representation is shown in Figure 11.

Table 17. Partial pressures of species in a series reaction.

Time	TMAP7 [A]	Theory [A]	Variance [A]	TMAP7 [B]	Theory [B]	Variance [B]	TMAP7 [C]	Theory [C]	Variance [C]
0	1.00E-06	1.00E-06	-0.00014	0.00E+00	0.00E+00	0.00000	0.00E+00	0.00E+00	0.00000
1	9.87E-07	9.88E-07	-0.00014	1.24E-08	1.24E-08	-0.00021	1.57E-11	1.55E-11	0.00977
2	9.75E-07	9.75E-07	-0.00014	2.46E-08	2.46E-08	-0.00021	6.22E-11	6.19E-11	0.00477
3	9.63E-07	9.63E-07	-0.00014	3.67E-08	3.67E-08	-0.00022	1.39E-10	1.39E-10	0.00313
4	9.51E-07	9.51E-07	-0.00015	4.85E-08	4.85E-08	-0.00020	2.46E-10	2.45E-10	0.00227
5	9.39E-07	9.39E-07	-0.00014	6.02E-08	6.02E-08	-0.00020	3.82E-10	3.81E-10	0.00177
6	9.28E-07	9.28E-07	-0.00014	7.17E-08	7.17E-08	-0.00022	5.47E-10	5.46E-10	0.00143
7	9.16E-07	9.16E-07	-0.00014	8.30E-08	8.30E-08	-0.00023	7.40E-10	7.39E-10	0.00117
8	9.05E-07	9.05E-07	-0.00014	9.42E-08	9.42E-08	-0.00023	9.62E-10	9.61E-10	0.00099
9	8.93E-07	8.94E-07	-0.00013	1.05E-07	1.05E-07	-0.00021	1.21E-09	1.21E-09	0.00083
10	8.82E-07	8.82E-07	-0.00013	1.16E-07	1.16E-07	-0.00023	1.49E-09	1.49E-09	0.00072
11	8.71E-07	8.72E-07	-0.00013	1.27E-07	1.27E-07	-0.00020	1.79E-09	1.79E-09	0.00060
12	8.61E-07	8.61E-07	-0.00013	1.37E-07	1.37E-07	-0.00023	2.12E-09	2.12E-09	0.00053
13	8.50E-07	8.50E-07	-0.00012	1.47E-07	1.48E-07	-0.00026	2.48E-09	2.48E-09	0.00046
14	8.39E-07	8.39E-07	-0.00013	1.58E-07	1.58E-07	-0.00023	2.86E-09	2.86E-09	0.00039
15	8.29E-07	8.29E-07	-0.00013	1.68E-07	1.68E-07	-0.00022	3.27E-09	3.26E-09	0.00035
16	8.19E-07	8.19E-07	-0.00012	1.78E-07	1.78E-07	-0.00024	3.70E-09	3.70E-09	0.00030
17	8.08E-07	8.09E-07	-0.00012	1.87E-07	1.87E-07	-0.00025	4.15E-09	4.15E-09	0.00027
18	7.98E-07	7.99E-07	-0.00012	1.97E-07	1.97E-07	-0.00026	4.63E-09	4.63E-09	0.00024
19	7.89E-07	7.89E-07	-0.00012	2.06E-07	2.06E-07	-0.00023	5.14E-09	5.14E-09	0.00025
20	7.79E-07	7.79E-07	-0.00013	2.15E-07	2.16E-07	-0.00021	5.66E-09	5.66E-09	0.00023
21	7.69E-07	7.69E-07	-0.00013	2.25E-07	2.25E-07	-0.00022	6.22E-09	6.21E-09	0.00023
22	7.59E-07	7.60E-07	-0.00013	2.34E-07	2.34E-07	-0.00022	6.79E-09	6.79E-09	0.00024
23	7.50E-07	7.50E-07	-0.00013	2.42E-07	2.42E-07	-0.00021	7.38E-09	7.38E-09	0.00023
24	7.41E-07	7.41E-07	-0.00013	2.51E-07	2.51E-07	-0.00017	8.00E-09	8.00E-09	0.00023
25	7.32E-07	7.32E-07	-0.00013	2.60E-07	2.60E-07	-0.00018	8.64E-09	8.64E-09	0.00022
26	7.22E-07	7.23E-07	-0.00013	2.68E-07	2.68E-07	-0.00017	9.30E-09	9.30E-09	0.00022
27	7.13E-07	7.14E-07	-0.00014	2.76E-07	2.76E-07	-0.00018	9.98E-09	9.98E-09	0.00022
28	7.05E-07	7.05E-07	-0.00014	2.85E-07	2.85E-07	-0.00018	1.07E-08	1.07E-08	0.00021
29	6.96E-07	6.96E-07	-0.00015	2.93E-07	2.93E-07	-0.00015	1.14E-08	1.14E-08	0.00023
30	6.87E-07	6.87E-07	-0.00014	3.01E-07	3.01E-07	-0.00016	1.21E-08	1.21E-08	0.00025

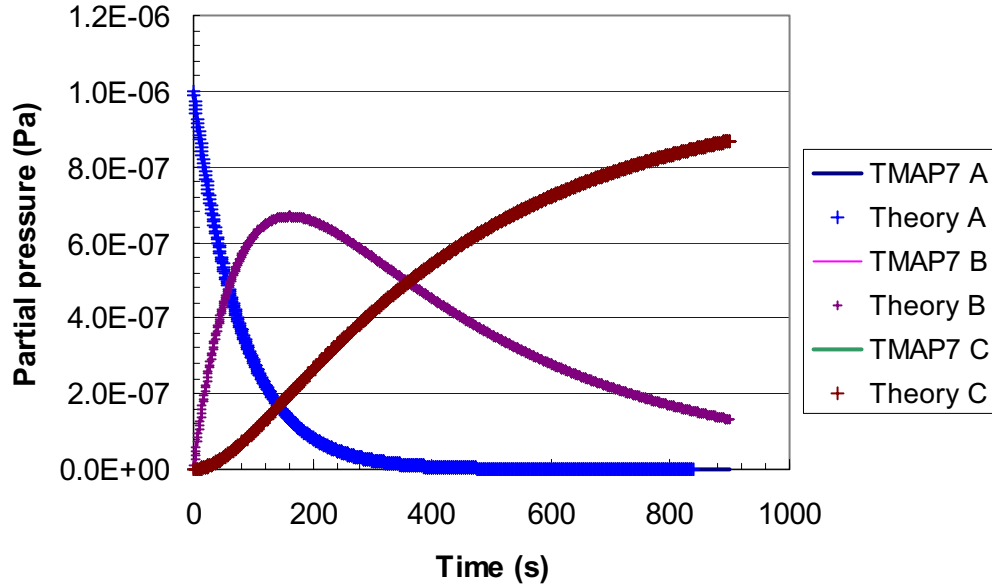


Figure 11. Partial pressures of species in series reaction (Val-1gc).

2.8 Problem 1h: Flow Through Multiple Enclosures

These two problems are designed to model convective flow between enclosures. The first problem models three enclosures. The first enclosure is a *boundary* enclosure whose concentration is constant. A convective flow goes from enclosure 1, through enclosure 2, to enclosure 3, and then back to enclosure 1. In the second problem, two enclosures are pre-charged with different species and a convective flow is allowed to circulate the species between the two enclosures.

2.8.1 Three Enclosure Problem (Val-1ha)

A system of three enclosures is modeled with flow from 1, to 2, to 3, and back to 1. Since enclosure 1 is defined as a boundary enclosure, concentration is constant. This enclosure acts as a source and a sink. The flux, \bar{j}_i , of molecules entering into enclosure i is given by

$$\bar{j}_i = QC_{i-1} \quad (42)$$

where

Q = volumetric flow rate, common for all enclosures ($0.1 \text{ m}^3/\text{sec}$)

C_{i-1} = concentration of gas molecules in enclosure $i-1$.

As the gas flows through the system, the number of atoms of the species of interest entering the 2nd and 3rd enclosures is greater than the number exiting. The concentration of that species in the enclosures rises towards the concentration in enclosure 1. The rate of change of the concentration of this species in the 2nd and 3rd enclosures can be modeled as follows

$$\begin{aligned}\frac{\partial C_2}{\partial t} &= \frac{Q(C_1 - C_2)}{V_2} \\ \frac{\partial C_3}{\partial t} &= \frac{Q(C_2 - C_3)}{V_3}\end{aligned}\tag{43}$$

The solution of this set of simultaneous equations with the initial condition that $C_2 = C_3 = 0$ is

$$C_2 = C_1 \left[1 - \exp\left(-\frac{Qt}{V_2}\right) \right]\tag{44}$$

and, if $V_2 = V_3$,

$$C_3 = C_1 \left[1 - \left(1 + \frac{Qt}{V_2} \right) \exp\left(-\frac{Qt}{V_2}\right) \right]\tag{45}$$

Otherwise C_3 is given by

$$C_3 = C_1 \left[1 - \frac{V_2}{V_2 - V_3} \exp\left(-\frac{Qt}{V_2}\right) + \frac{V_3}{V_2 - V_3} \exp\left(-\frac{Qt}{V_3}\right) \right]\tag{46}$$

In this problem, the following values were used to solve Eqs. (44) and (45),

$$\begin{aligned}V_2 &= V_3 = 1 \text{ m}^3, \\ C_1 &= 2.3904 \text{ atoms/m}^3, \\ Q &= 0.1 \text{ m}^3/\text{sec}.\end{aligned}$$

The values of Eqs. (44) and (45), were converted to partial pressures and solved using MS Excel and are compared with the values obtained from TMAP7 in Table 18 and Figure 12. The variance for Enclosure 2 is less than 0.1% for all time, while the variance for Enclosure 3 is always below 0.3%, and both seem to be random with time.

2.8.2 Equilibrating Enclosures (Val-1hb)

The second flow problem is setup as a system of two enclosures, each with a volume of 1 m^3 , with flow from enclosures 1 to 2, and 2 to 1. Enclosure 1 is initially charged with tritium (T) and enclosure 2 is pre-charged with deuterium (D), each at 1 Pa. The concentration change rates for this system are given by the following for species T.

$$\begin{aligned}\frac{dC_{T_1}}{dt} &= \frac{Q}{V} (C_{T_2} - C_{T_1}) \\ \frac{dC_{T_2}}{dt} &= \frac{Q}{V} (C_{T_1} - C_{T_2})\end{aligned}\tag{47}$$

and for species D

Table 18. Concentration profiles of enclosures 2 and 3 with convective flow.

Time (s)	Enclosure 2			Enclosure 3		
	TMAP7	Theory	Var 1	TMAP7	Theory	Var 1
0	0.00E+00	0.00E+00	0	0.00E+00	0.00E+00	0
1	9.51E-02	9.52E-02	-0.00021	4.68E-03	4.68E-03	0.00070
2	1.81E-01	1.81E-01	-0.00011	1.75E-02	1.75E-02	0.00039
3	2.59E-01	2.59E-01	-0.00012	3.69E-02	3.69E-02	0.00018
4	3.30E-01	3.30E-01	-0.00030	6.15E-02	6.16E-02	-0.00026
5	3.93E-01	3.93E-01	-0.00038	9.02E-02	9.02E-02	-0.00049
6	4.51E-01	4.51E-01	-0.00042	1.22E-01	1.22E-01	-0.00067
7	5.03E-01	5.03E-01	-0.00045	1.56E-01	1.56E-01	-0.00074
8	5.51E-01	5.51E-01	-0.00026	1.91E-01	1.91E-01	-0.00030
9	5.93E-01	5.93E-01	-0.00007	2.28E-01	2.28E-01	0.00010
10	6.32E-01	6.32E-01	0.00005	2.64E-01	2.64E-01	0.00041
11	6.67E-01	6.67E-01	0.00015	3.01E-01	3.01E-01	0.00063
12	6.99E-01	6.99E-01	0.00021	3.38E-01	3.37E-01	0.00079
13	7.28E-01	7.27E-01	0.00025	3.74E-01	3.73E-01	0.00089
14	7.54E-01	7.53E-01	0.00027	4.09E-01	4.08E-01	0.00099
15	7.77E-01	7.77E-01	0.00030	4.43E-01	4.42E-01	0.00105
16	7.98E-01	7.98E-01	0.00006	4.75E-01	4.75E-01	0.00042
17	8.17E-01	8.17E-01	-0.00015	5.07E-01	5.07E-01	-0.00015
18	8.34E-01	8.35E-01	-0.00031	5.37E-01	5.37E-01	-0.00058
19	8.50E-01	8.50E-01	-0.00044	5.66E-01	5.66E-01	-0.00096
20	8.64E-01	8.65E-01	-0.00054	5.93E-01	5.94E-01	-0.00125
21	8.77E-01	8.78E-01	-0.00061	6.19E-01	6.20E-01	-0.00149
22	8.89E-01	8.89E-01	-0.00066	6.44E-01	6.45E-01	-0.00169
23	8.99E-01	9.00E-01	-0.00070	6.68E-01	6.69E-01	-0.00183
24	9.09E-01	9.09E-01	-0.00072	6.90E-01	6.92E-01	-0.00194
25	9.17E-01	9.18E-01	-0.00074	7.11E-01	7.13E-01	-0.00202
26	9.25E-01	9.26E-01	-0.00060	7.31E-01	7.33E-01	-0.00160
27	9.32E-01	9.33E-01	-0.00036	7.51E-01	7.51E-01	-0.00084
28	9.39E-01	9.39E-01	-0.00016	7.69E-01	7.69E-01	-0.00018
29	9.45E-01	9.45E-01	0.00000	7.86E-01	7.85E-01	0.00037
30	9.50E-01	9.50E-01	0.00012	8.02E-01	8.01E-01	0.00083
31	9.55E-01	9.55E-01	0.00023	8.16E-01	8.15E-01	0.00120
32	9.60E-01	9.59E-01	0.00030	8.30E-01	8.29E-01	0.00152
33	9.63E-01	9.63E-01	0.00036	8.43E-01	8.41E-01	0.00178
34	9.67E-01	9.67E-01	0.00041	8.55E-01	8.53E-01	0.00197
35	9.70E-01	9.70E-01	0.00043	8.66E-01	8.64E-01	0.00213
36	9.73E-01	9.73E-01	0.00045	8.76E-01	8.74E-01	0.00224
37	9.76E-01	9.75E-01	0.00036	8.86E-01	8.84E-01	0.00192
38	9.78E-01	9.78E-01	0.00010	8.93E-01	8.93E-01	0.00091
39	9.80E-01	9.80E-01	-0.00010	9.01E-01	9.01E-01	0.00004
40	9.81E-01	9.82E-01	-0.00028	9.08E-01	9.08E-01	-0.00070

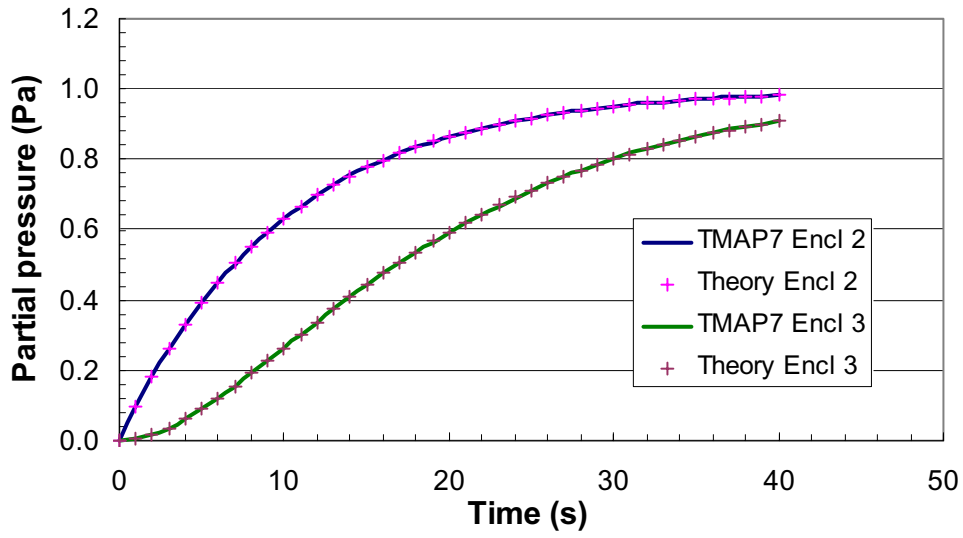


Figure 12. Concentration history of sequentially coupled enclosures (Val-1ha).

$$\begin{aligned}\frac{dC_{D_1}}{dt} &= \frac{Q}{V}(C_{D_2} - C_{D_1}) \\ \frac{dC_{D_2}}{dt} &= \frac{Q}{V}(C_{D_1} - C_{D_2})\end{aligned}\tag{48}$$

where

Q = volumetric flow ($0.1 \text{ m}^3/\text{s}$)

V = volume (1 m^3)

C_{T_i} = concentration of tritium in Enclosure i

C_{D_i} = concentration of deuterium in Enclosure i

A mass balance on the system, gives a relationship between the concentration of species in Enclosure 1 and Enclosure 2.

$$C_{T_1} = C_{T_1}^o - C_{T_2}\tag{49}$$

Now by substituting Eq. (49) into the first of Eqs. (48), the solution is given by

$$C_{T_1} = \frac{C_{T_1}^o}{2} \left[1 + \exp\left(-\frac{2Q}{V}t\right) \right]\tag{50}$$

where $C_{T_1}^o$ = initial concentration of tritium in Enclosure 1.

It is recognized that for the same initial starting conditions for deuterium, except different initial pressures (1 Pa in enclosure 2 and 0 Pa in enclosure 1), the following will be true

$$\begin{aligned} C_{D_2} &= C_{T_1} \\ C_{D_1} &= C_{T_2} \end{aligned} \quad (51)$$

Eq. (50) was solved in Excel™ and compared with the values obtained from TMAP7. These values, presented as partial pressures, are listed in Table 19 and shown graphically in Figure 13.

Table 19. Concentration of tritium in convective flow between two enclosures

Time (s)	Enclosure 1			Enclosure 2		
	TMAP7	Theory	Variance	TMAP7	Theory	Variance
0	1.000	1.000	0.00014	0.000	0.000	0.00000
1	0.909	0.909	0.00014	0.091	0.091	0.00024
2	0.835	0.835	0.00014	0.165	0.165	0.00012
3	0.774	0.774	0.00012	0.226	0.226	0.00024
4	0.725	0.725	0.00008	0.275	0.275	0.00035
5	0.684	0.684	0.00003	0.316	0.316	0.00038
6	0.651	0.651	0.00006	0.349	0.349	0.00029
7	0.623	0.623	0.00016	0.377	0.377	0.00011
8	0.601	0.601	0.00025	0.399	0.399	0.00000
9	0.582	0.583	0.00029	0.417	0.417	-0.00007
10	0.567	0.568	0.00033	0.432	0.432	-0.00009
11	0.555	0.555	0.00022	0.445	0.445	0.00006
12	0.545	0.545	0.00007	0.455	0.455	0.00024
13	0.537	0.537	-0.00002	0.463	0.463	0.00035
14	0.530	0.530	-0.00010	0.469	0.470	0.00042
15	0.525	0.525	-0.00013	0.475	0.475	0.00046
16	0.520	0.520	-0.00015	0.479	0.480	0.00046
17	0.517	0.517	0.00007	0.483	0.483	0.00021
18	0.514	0.514	0.00024	0.486	0.486	0.00006
19	0.511	0.511	0.00032	0.489	0.489	-0.00005
20	0.509	0.509	0.00039	0.491	0.491	-0.00012
21	0.507	0.507	0.00043	0.493	0.493	-0.00014
22	0.506	0.506	0.00037	0.494	0.494	-0.00008
23	0.505	0.505	0.00015	0.495	0.495	0.00015
24	0.504	0.504	0.00001	0.496	0.496	0.00029
25	0.503	0.503	-0.00010	0.496	0.497	0.00038
26	0.503	0.503	-0.00016	0.497	0.497	0.00045
27	0.502	0.502	-0.00018	0.498	0.498	0.00049
28	0.502	0.502	-0.00020	0.498	0.498	0.00048
29	0.502	0.502	-0.00001	0.498	0.498	0.00031
30	0.501	0.501	0.00016	0.499	0.499	0.00012

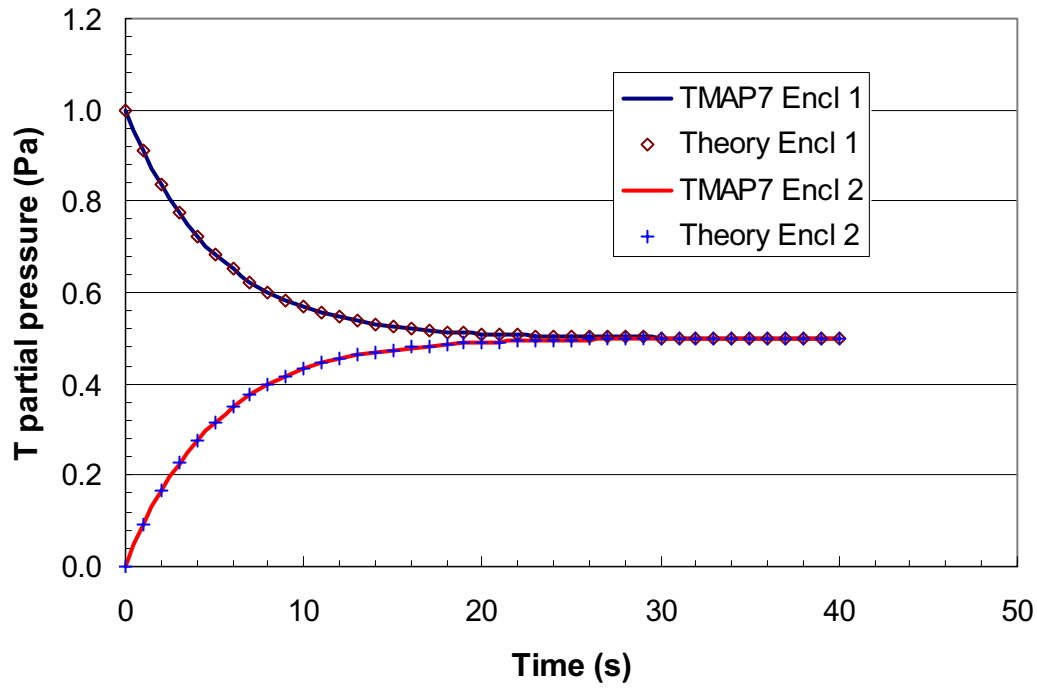


Figure 13. Tritium concentration equilibration in two communicating enclosures (Val-1hb).

2.9 Problem 1i: Species Equilibration on a Reactive Surface

When two species can react on a surface to form a third, it is possible to predict the rate at which equilibration between the species will occur. For example, consider the reaction between two isotopic species



2.9.1 Ratedep Conditions (Val-1ia, Val-1ib)

The expression (derived in Appendix A) for the rate of formation of AB when the conversion rate at the surface is high is

$$P_{AB} = \frac{2P_{A_2}^0 P_{B_2}^0}{P_{A_2}^0 + P_{B_2}^0} \left[1 - \exp\left(-\frac{SK_d kT}{V} t\right) \right] \quad (53)$$

Here

$P_{A_2}^o$ = initial partial pressure of species A_2 in the enclosure

$P_{B_2}^o$ = initial partial pressure of species B_2 in the enclosure

S = surface area available for reaction

K_d = dissociation coefficient of both A_2 and B_2 on the surface

k = Boltzmann's constant

T = temperature

V = enclosure volume

In this case A was taken as protium, and B was deuterium. S was assumed to be 0.0025 m^2 , K_d for $M = 2 \text{ amu}$ is $1.85804\text{E}+24 / \sqrt{T} \text{ atom/m}^2/\text{s}$, T was $1,000 \text{ K}$, and V was 1 m^3 . $P_{A_2}^o$ and $P_{B_2}^o$ were first assumed equal at $1.0 \times 10^4 \text{ Pa}$ and then $P_{B_2}^o$ was assumed to be increased to $1.0 \times 10^5 \text{ Pa}$. Figure 14 shows the comparison of TMAP7 with the theory for the case of equal starting partial pressures, and Figure 15 shows the comparison for the unequal starting partial pressure case. Variance was less than 1% for all times and generally less than 0.1%.

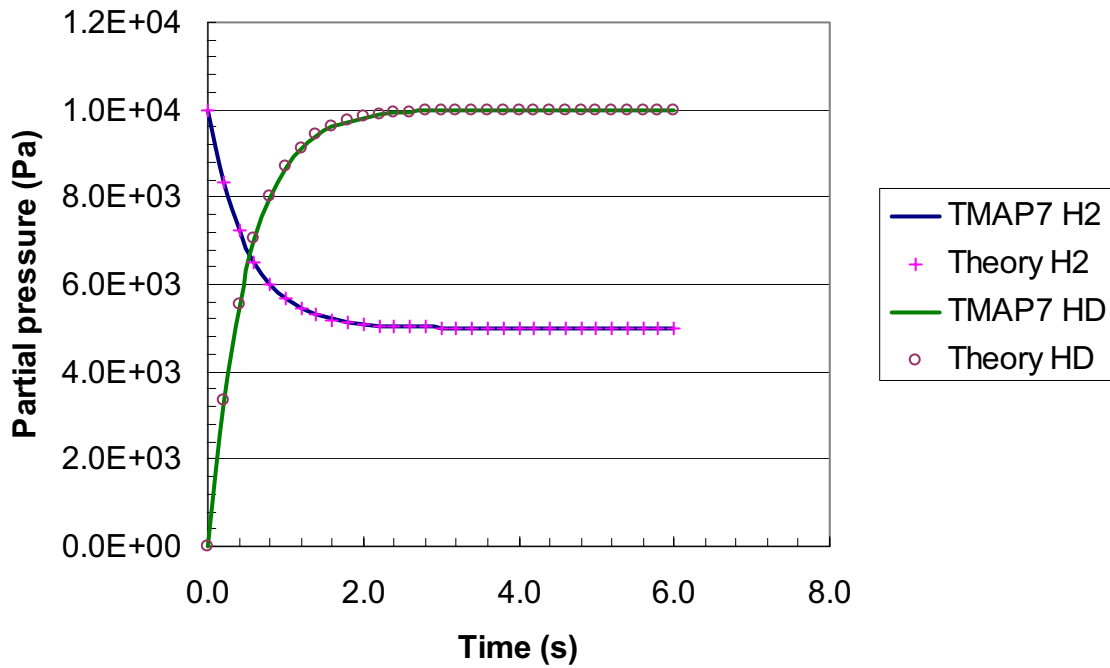


Figure 14. Equilibration of H_2 with D_2 to form HD on a tungsten surface under the assumption of equal starting partial pressures for the reactants and *ratedep* boundary conditions (Val-1ia).

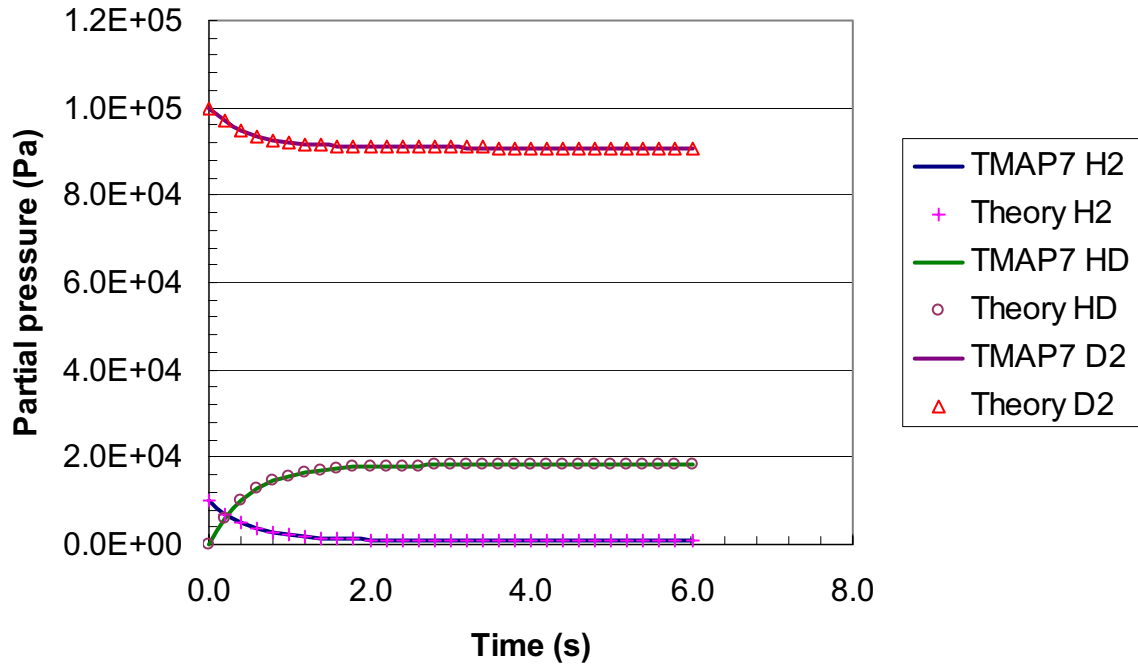


Figure 15. Chemical equilibration of H₂ and D₂ to HD on a tungsten surface with unequal starting partial pressures and *ratedep* boundary conditions (Val-1ib).

2.9.2 Surfdep Conditions (Val-1ic, Val-1id)

When surface processes are governed by activation energies with dissociation and recombination considered explicitly, *surfdep* boundary conditions govern. As explained in Appendix A, the equation for transient pressure of *AB* given starting pressures of *A*₂ and *B*₂ is

$$P_{AB} = \frac{2P_{A_2}^0 P_{B_2}^0}{P_{A_2}^0 + P_{B_2}^0} \left[1 - \exp\left(-\frac{t}{\tau}\right) \right] \quad (54)$$

Where

$$\tau = \frac{V(\hat{K}_r + K_b)}{SkT \hat{K}_d K_b} \quad (55)$$

and for molecular mass *M*,

$$K_d = \frac{1}{\sqrt{2\pi MkT}} \quad (56)$$

K_b is a thermally activated dissociation coefficient, assumed to be given by a Boltzmann equation with activation energy *E_b*, and Debye frequency *ν_o*.

$$K_b = \nu_o \exp\left(-\frac{E_b}{kT}\right) \quad (57)$$

Molecules escape from the surface at rate

$$\hat{K}_r = \frac{\nu_o}{6} \exp\left(\frac{E_c - E_x}{kT}\right) \quad (58)$$

Here, in addition to variables previously defined, E_c is the surface binding energy, and the factor of 6 accounts for the probability that on any given vibration, the direction of the phonon is away from the material surface.

The first of the *surfep* cases uses equal starting pressures of 1.0×10^4 Pa of H_2 and D_2 and no HD. In this case, E_x was specified to 0.05 eV, E_c was -0.01 eV and the dissociation energy was taken as zero, meaning that attempts at the Debye frequency all succeed. Temperature was again 1,000 K, the surface area was a 5-cm x 5-cm square, and the enclosure volume was 1.0 m^3 .

Comparison of TMAP7 code results with the theoretical values is made in Figure 16.

Corresponding results for unequal starting pressures are shown in Figure 17.

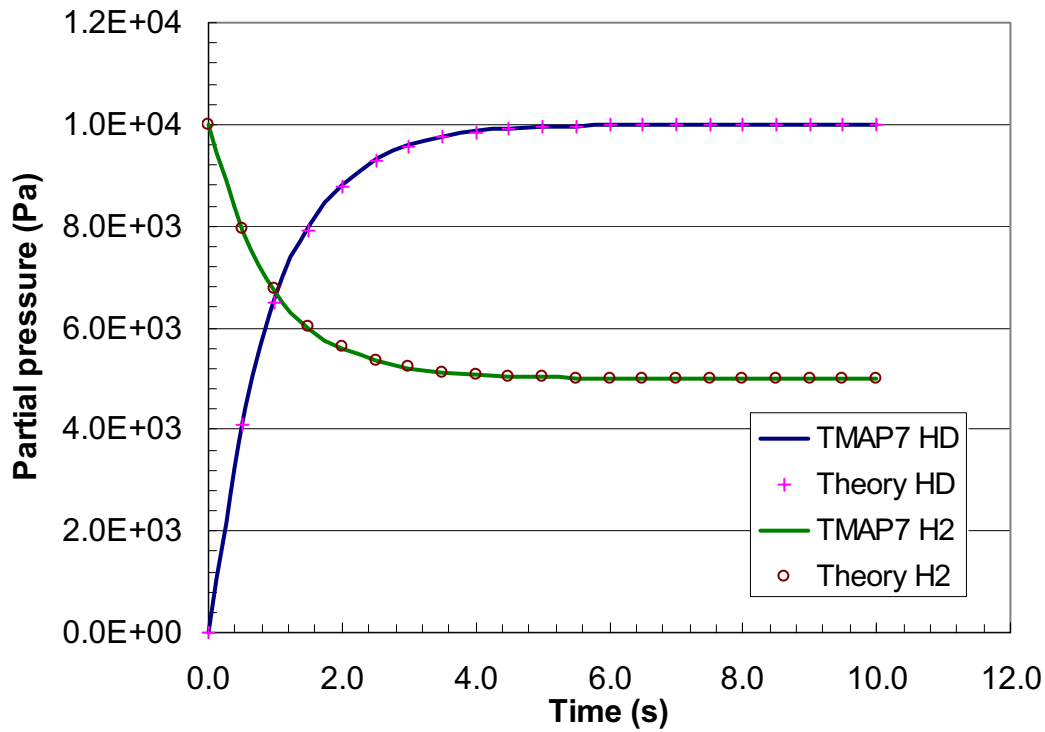


Figure 16. Chemical equilibration of H_2 and D_2 to form HD under *surfdep* boundary conditions with equal starting pressures (Val-1ic).

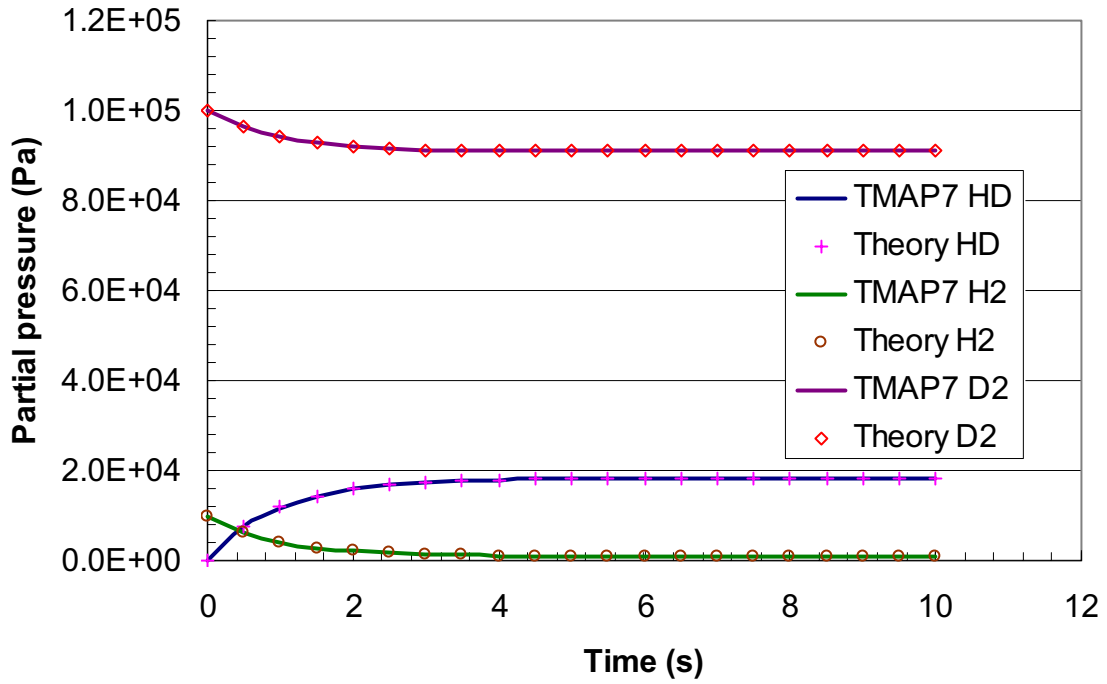


Figure 17. Chemical equilibration of H₂ and D₂ to form HD under *surfdep* boundary conditions with unequal starting pressures (Val-1id).

2.10 Problem 1j: Radioactive Decay

Two problems were run to demonstrate tritium decay, though any other isotope could have been chosen. The first is simple decay of mobile species in a slab. The second is decay of trapped atoms in a similar slab but with a distributed trap concentration.

2.10.1 Problem 1ja: Radioactive Decay of Mobile Tritium in a Slab (Val-1ja)

This model is employed to test the first order radioactive decay capabilities of TMAP7. The model assumes pre-charging of a slab with tritium. The tritium was uniformly distributed over the thickness of the slab. The tritium decays to ³He as shown in Equation (1) with a half-life of 12.3232 years.



The concentrations of the two species are calculated. The concentration of T at any given time is given by

$$C_t = C_t^o \exp(-\omega t) \quad (60)$$

Applying a mass balance over the system, the concentration of helium is given by

$$C_{{}^3\text{He}} = C_t^o [1 - \exp(-\omega t)] \quad (61)$$

where

C_t^o = Initial concentration of tritium

ω = rate constant ($1.78241\text{E-}9 \text{ s}^{-1}$)

t = time (sec).

The comparison between the TMAP7 result and Eqs. (60) and (61) for mobile tritium can be seen in Table 20. A graphical representation is given in Figure 18.

Table 20. Decay of mobile tritium to ^3He (Val-1ja)

Time (yr)	TMAP7	[T] Theory	Variance	TMAP7	[He] Theory	Variance
0.0	1.00000	1.00000	0.00000	0.00000	0.00000	0.00000
0.4	0.97973	0.97971	0.00002	0.02009	0.02029	-0.01001
0.7	0.95987	0.95983	0.00003	0.03997	0.04017	-0.00499
1.1	0.94033	0.94036	-0.00003	0.05944	0.05964	-0.00333
1.5	0.92127	0.92128	-0.00001	0.07852	0.07872	-0.00254
1.8	0.90260	0.90259	0.00001	0.09722	0.09741	-0.00197
2.2	0.88427	0.88428	-0.00001	0.11553	0.11572	-0.00165
2.6	0.86633	0.86633	0.00000	0.13347	0.13367	-0.00144
2.9	0.84880	0.84876	0.00005	0.15105	0.15124	-0.00125
3.3	0.83153	0.83154	0.00000	0.16828	0.16846	-0.00109
3.6	0.81467	0.81467	0.00000	0.18515	0.18533	-0.00098
4.0	0.79813	0.79814	0.00000	0.20168	0.20186	-0.00091
4.4	0.78193	0.78194	-0.00001	0.21787	0.21806	-0.00084
4.7	0.76607	0.76608	-0.00002	0.23375	0.23392	-0.00075
5.1	0.75053	0.75054	0.00000	0.24929	0.24946	-0.00071
5.5	0.73533	0.73531	0.00003	0.26452	0.26469	-0.00065
5.8	0.72040	0.72039	0.00001	0.27944	0.27961	-0.00061
6.2	0.70580	0.70577	0.00004	0.29406	0.29423	-0.00057
6.6	0.69147	0.69145	0.00002	0.30838	0.30855	-0.00054
6.9	0.67747	0.67742	0.00006	0.32241	0.32258	-0.00052
7.3	0.66371	0.66368	0.00004	0.33615	0.33632	-0.00049
7.7	0.65025	0.65022	0.00005	0.34962	0.34978	-0.00047
8.0	0.63705	0.63702	0.00005	0.36282	0.36298	-0.00043
8.4	0.62413	0.62410	0.00005	0.37575	0.37590	-0.00041
8.7	0.61147	0.61144	0.00005	0.38841	0.38856	-0.00041
9.1	0.59906	0.59903	0.00005	0.40081	0.40097	-0.00039
9.5	0.58691	0.58688	0.00005	0.41297	0.41312	-0.00036
9.8	0.57500	0.57497	0.00005	0.42488	0.42503	-0.00035
10.2	0.56334	0.56330	0.00006	0.43655	0.43670	-0.00034
10.6	0.55191	0.55187	0.00006	0.44798	0.44813	-0.00032
10.9	0.54071	0.54068	0.00007	0.45918	0.45932	-0.00031
11.3	0.52974	0.52971	0.00006	0.47015	0.47029	-0.00031
11.7	0.51899	0.51896	0.00006	0.48090	0.48104	-0.00029
12.0	0.50847	0.50843	0.00007	0.49143	0.49157	-0.00029
12.4	0.49815	0.49812	0.00007	0.50175	0.50188	-0.00027

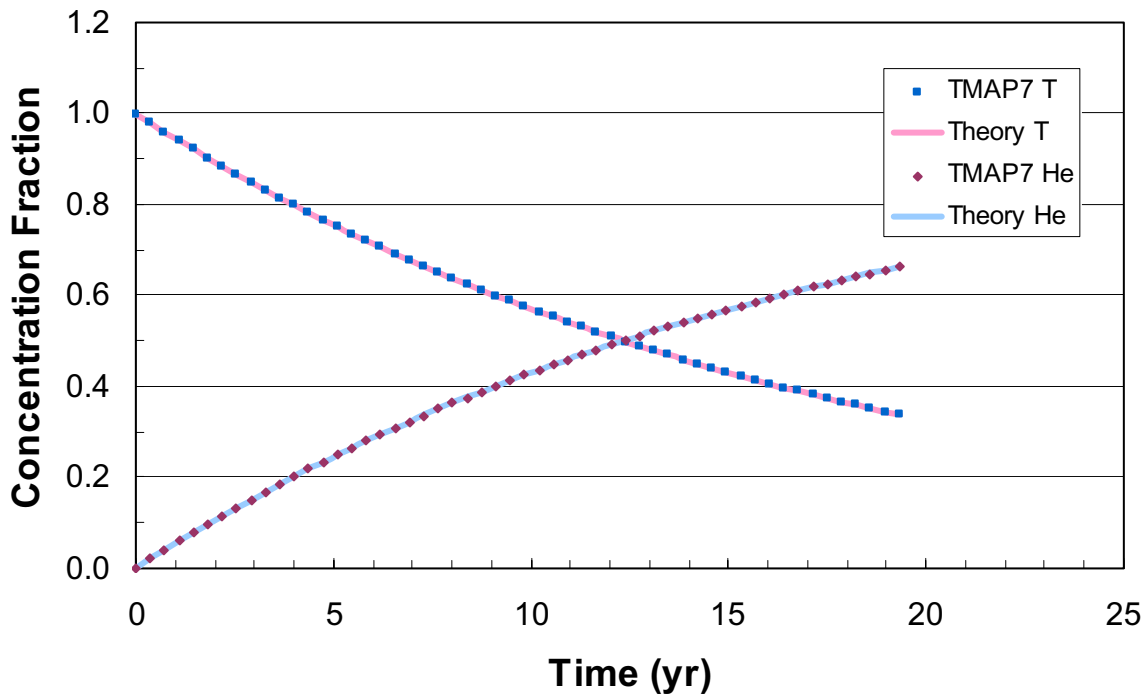


Figure 18. Decay of mobile tritium and associated growth of ^3He in a diffusion segment (Val-1ja).

2.10.2 Problem 1jb: Decay of Tritium in a Distributed Trap (Val-1jb)

A further but more complex exercise was run for a slab in which nearly all of the tritium is trapped. A slab similar to that used in Problem 1ja was used here, but traps at 1% atom fraction and 4.2-eV trap energy were distributed in a normal distribution centered at the mid-plane of the slab. The traps were initially filled to 10% of trap concentration. The mobile atom concentration was only 1 atom/m^3 to begin with, and it very quickly was all absorbed into these deep traps. This problem also demonstrates the utility of the pre-programmed distribution functions for certain parameters.

Figure 19 shows the depth profiles of initial trapped atoms of tritium, final trapped atoms of tritium after 45 years, and the distribution of He-3 at the end of that time. Note that because of finite diffusivity of the He-3, it has broadened a little from the trap concentration. The theoretical solution for this broadening is very complex and is not presented here.

Figure 20 shows the total inventory of tritium in the trap as a function of time over the decay period. It also shows the total helium inventory (atoms/m^2). The same precision as demonstrated in Problem 1ja was observed here.

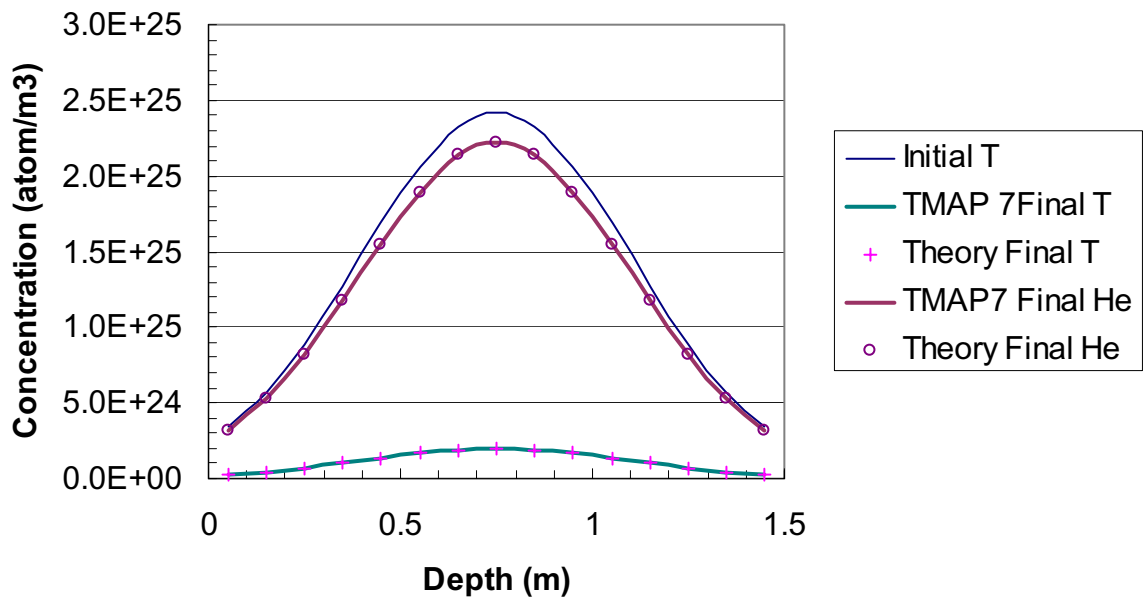


Figure 19. Profiles of trapped tritium at the beginning and end of a 45-year decay and the profile of the resultant ^3He at the end of that time (Val-1jb).

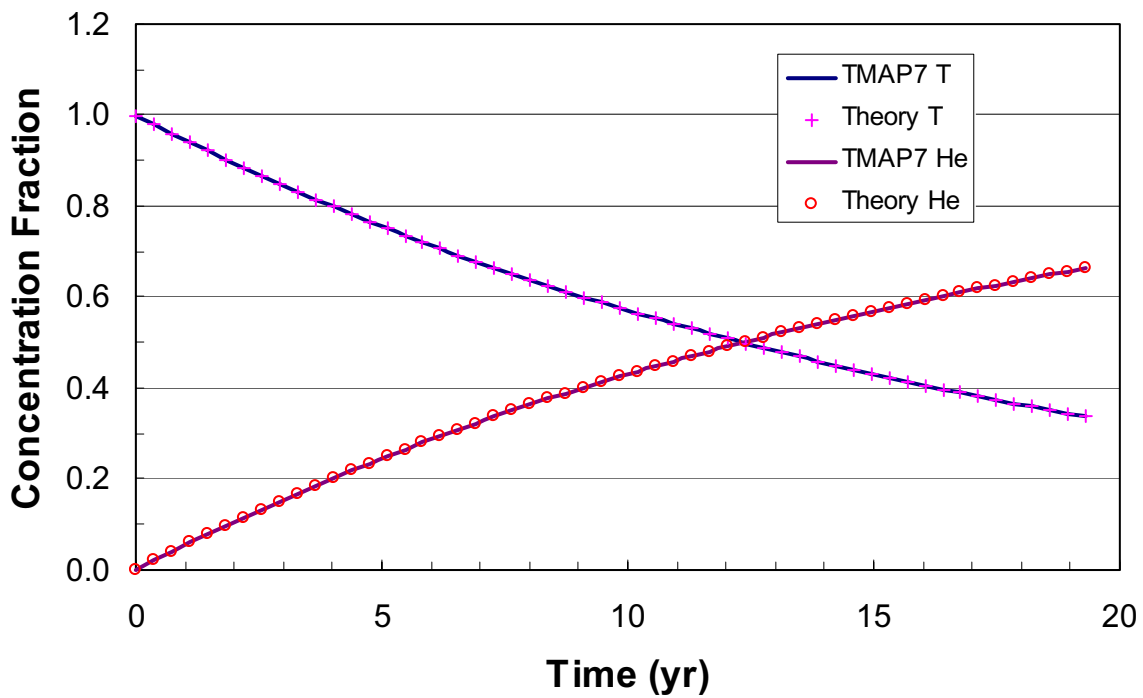


Figure 20. Loss of trapped tritium by radioactive decay is reflected in the gain of He-3 (Val-1jb).

3.0 REPLICATING EXPERIMENTS

The second phase of code validation is the comparison of code results with actual experimental data. Published experiments together with their experimental data were selected for modeling. The first three of these are repeats from the verification and validation of TMAP4.⁶

3.1 Problem 2a: Ion Implantation Experiment (Val-2a)

This problem is the simulation of experimental results obtained at the INEL in 1985 and published.¹⁴ The experiment involved applying an ion beam to a 2.5-cm diameter, 0.5-mm thick sample of a modified 316 stainless steel called Primary Candidate Alloy (PCA). Details of the experiment and the means of evaluating the necessary transport parameters to get a good fit between TMAP7 results and the experimental data are given in the publication. The TRIM code was used to determine that the average implantation depth for the ions was $11\text{-}\mu\text{m} \pm 5.4\text{ }\mu\text{m}$. Reemission data from the TRIM calculation showed that only 75% of the incident flux remained in the metal. The other 25% was re-emitted.

One known non-physical feature in the modeling is that the cleanup of the upstream surface was modeled by a simple exponential in time rather than an ion fluence which was interrupted twice during the actual experiment. The pressures upstream and downstream proved to be inconsequential; they could have been taken as zero and obtained essentially the same results.

The plot of Figure 15 was generated. Actual experimental data are also shown on the figure. They are fairly closely approximated by the calculated permeation. Notice in the figure, however, that in the experimental data there is a lower permeation flux value when the beam is on, and a relatively slow trail-off, compared with the calculation, when the beam was turned off. Some of this is a consequence of the experimental technique where the walls of the experimental chamber did some pumping of the gas as it came through the sample and then provided a source of deuterium when the sample permeation ceased. Some two-dimensional effects also influence the comparison.

Results of this calculation using TMAP7 are essentially identical to those obtained using TMAP4 and reported previously.

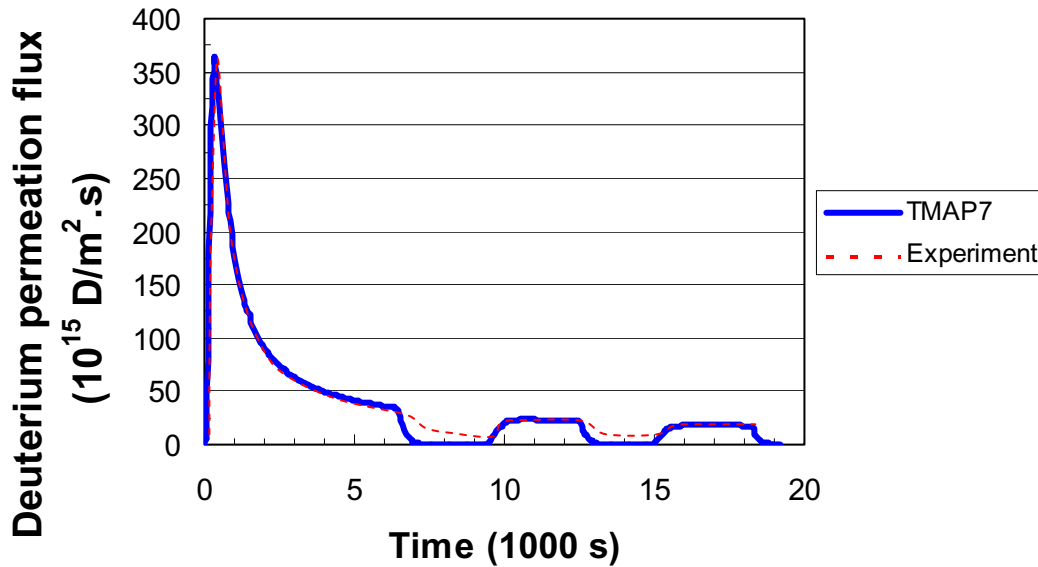


Figure 21. Plasma Driven Permeation of PCA (Val-2a)

3.2 Problem 2b: Diffusion Experiment in Beryllium (Val-2ba, Val-2bb)

This problem is taken from work done by R. G. Macaulay-Newcombe at McMaster University.¹⁵ He and co-workers conducted thermal absorption and desorption experiments, as well as implantation experiments, on wafers of polished beryllium. Of the several data sets presented, the one modeled here is that represented in Figure 12 (a) in their publication. The beryllium was 0.4-mm thick and had an area of 104 mm². It was polished to a mirror finish and then exposed to 13.3 kPa of deuterium at 773 K for 50 minutes. It was quickly cooled under a vacuum of about 1 μ Pa. The cooling time constant for the apparatus is taken as 45 minutes. After removing the sample from the charging furnace, it was transferred in the air to a thermal desorption furnace where the temperature was increased from ambient (300 K) to 1073 K at the rate of 3K/min. This was done under vacuum, and the pressure of the chamber was monitored by residual gas analysis and calibrated against standard leaks. In that way, the emission rate from the sample could be measured as a function of temperature. Data from that measurement, given in Figure 12 (a) of their paper are reproduced in Figure 22 here. From Rutherford backscattering measurements made on the samples before charging with deuterium, they deduced that the thickness of the oxide film was 18 nm. This is typical for polished beryllium. The metal is so reactive in air that the film forms almost immediately after any surface oxide removal. On the other hand, it is relatively stable and would only grow slightly when exposed to air between charging and thermal desorption.

This experiment is modeled using a two-segment model in TMAP7 with the segments linked. The first is the BeO film, which is modeled using equally spaced nodes of 1 nm each plus the two surface nodes. The second segment is a half-thickness wafer of beryllium with reflective boundary conditions at the mid-plane. It is made up of 15 segments of varying thickness to

accommodate solution stiffness plus the two surface nodes. The solubility of deuterium in beryllium used was that given by K. L. Wilson, et al.,¹⁶ based on work done by W. A. Swansiger, also of Sandia National Laboratory. The diffusivity of deuterium in beryllium was measured by E. Abramov, et. al.¹⁷. They made measurements on high-grade (99% pure) and extra-grade (99.8% pure). The values used here are those for high-grade beryllium, consistent with Dr. Macaulay-Newcombe's measurements of the purity of his samples.

Deuterium transport properties of the BeO are more challenging. First, it is not clear in what state the deuterium exists in the BeO. However, it has been observed¹⁸ that an activation energy of -78 KJ.mole (exothermic solution) is evident for tritium coming out of neutron irradiated beryllium in work done by D. L. Baldwin of Battelle Pacific Northwest Laboratory. The same energy has appeared in other results (can be inferred from Dr. Swansiger's work cited by Wilson, et al.¹⁶, and by R. A. Causey, et al.¹⁹, among others), so one may be justified in using it. The solubility coefficient is not well known. Measurements reported by R. G. Macaulay-Newcombe, et al.²⁰ and in follow-up conversations indicate about 200 appm of D in BeO after exposure to 13.3 kPa of D₂ at 773 K. That suggests a coefficient of only $1.88 \times 10^{-18} \text{ d/m}^3 \text{Pa}^{1/2}$. Since much of the deuterium in the oxide layer will get out during the cool-down process (and because it gives a good fit) the solubility coefficient is taken to be $5 \times 10^{-20} \text{ d/m}^3 \text{Pa}^{1/2}$.

Deuterium diffusion measurements in BeO were made by J. D. Fowler, et al.²¹. They found a wide range of results for diffusivity in BeO, depending on the physical form of the material, having measured it for single-crystal, sintered, and powdered BeO. This model uses one expression for the charging phase and another for the thermal desorption phase, believing that the surface film changed somewhat during the transfer between the two furnaces. For the charging phase diffusivity, the model uses 20 times that for the sintered BeO. Thermal expansion mismatches tend to open up cracks and channels in the oxide layer, so this seems a reasonable value. The same activation energy of 48.5 kJ/mole, is retained, however. For the thermal desorption phase, the diffusivity prefactor of the sintered material ($7 \times 10^{-5} \text{ m}^2/\text{sec}$) and an activation energy of 223.7 kJ/mole (53.45 kcal/mole) are used. These values give good results and lie well within the scatter of Fowlers data. Exposure of the sample to air after heating should have made the oxide more like single crystal by healing the cracks that may have developed.

The model applies 13.3. kPa of D₂ for 50 hours followed by evacuation to 1 μPa and cool down with a 45 minute time constant for one hour. The deuterium concentrations in the sample are of a complex distribution that results from first charging the sample and then discharging it during the cool down. This problem is then restarted with different equations to simulate thermal desorption in the 1- μPa environment. That begins at 300 K and goes to 1073 K. Again, the concentration profiles in both the substrate beryllium and the oxide film have a peculiar interaction because of the activation energies involved, but the flux exuding from the sample when doubled to account for the two sides of the specimen in the laboratory gives a good fit to the experimental data.

From the extracted diffusion species surface flux data for the left side of thermseg/diffseg 1, the solid curve in Figure 22 is constructed where it is compared with the experimental data. Agreement is virtually identical with that found in the TMAP4 calculation for this problem⁶.

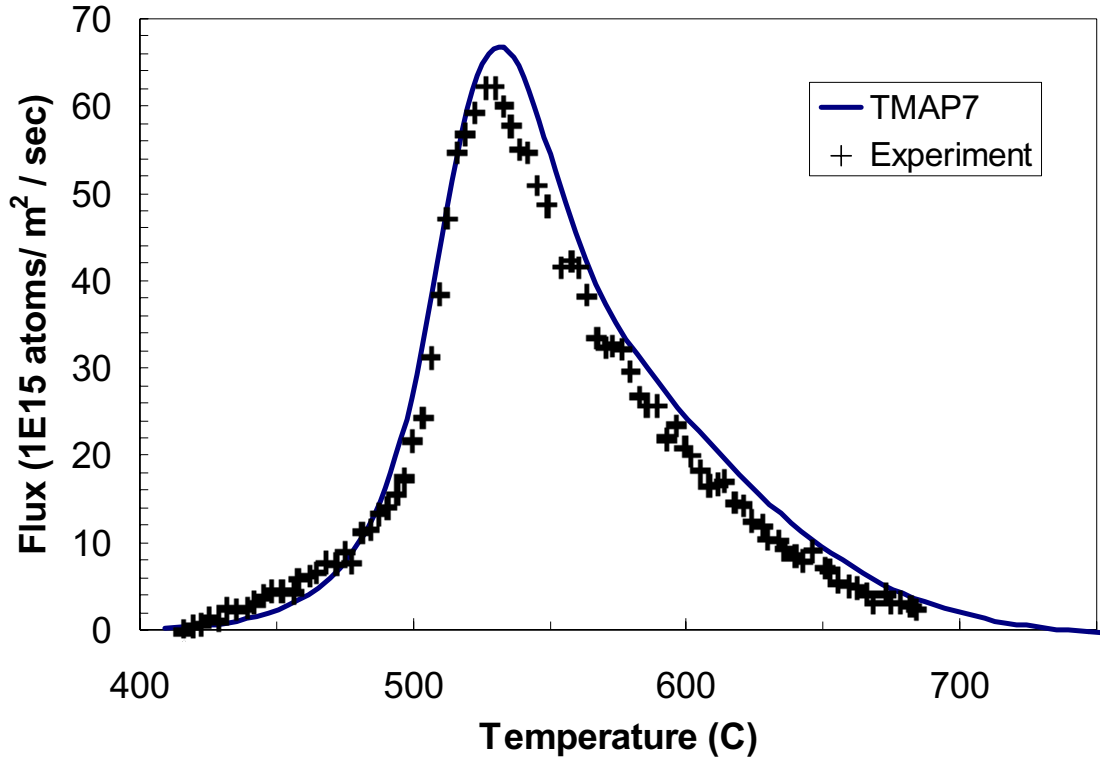


Figure 22. Thermal desorption test of beryllium (Val-2b)

3.3 Problem 2c: Test Cell Release Experiment (Val-2c)

This is an experiment that involves multiple enclosures and chemical reactions. It was conducted at the Tritium Systems Test Assembly (TSTA) at Los Alamos National Laboratory and documented by Holland and Jalbert.²² The main part of the experiment was an exposure chamber with a nominal volume of 1 m³, which was lined with epoxy paint that is 0.16 mm thick. Tritium was admitted to the chamber as T₂ at the commencement of the experiment. Normally moist (20% R.H.) air was admitted to the chamber at the rate of 0.54m³/hr constantly throughout the test. Samples of glycol taken from a bubbler just downstream from the exposure chamber were taken at intervals and scintillation counted to determine the time averaged HTO concentration in the chamber as a histogram in time. Tritium and water were absorbed into the paint during the initial part of the test and re-emitted later. Chemical reactions described by the formulae



took place within the exposure chamber, mainly as a consequence of the radioactivity of the tritium itself. Results of Holland and Jalbert are shown in their Figure 3 from the measurements of the resulting HTO concentration in the exposure chamber following a 10 Ci initial injection (effectively instantaneously) while purging with room air.

The TMAP7 Model for this experiment consists of three enclosures (1) the room from which air is drawn, (2) the exposure chamber, and (3) the tritium waste treatment system (TWT) to which the exhaust gases are directed. Only enclosure (2) is treated as "functional" or chemically active. The paint on the inside of the exposure chamber is treated as a diffusive segment and non-flow conditions are employed at the interface of the paint with the underlying aluminum foil. Experiments had previously demonstrated that there is virtually no transport of tritium into the aluminum foil. The techniques for determining the constants and other information required to generate a model that gives reasonable results are given by Holland and Jalbert and are not duplicated here.

Data were calculated by TMAP7 for the HTO concentration in the exposure chamber, enclosure 2. A pumping rate of $0.43 \text{ m}^3/\text{hr}$ gave a better fit than the apparent one of $0.54 \text{ m}^3/\text{hr}$. A solid curve representing these data is compared in Figure 23 with measurements made in bubblers in line with the exposure chamber exhaust. The period over which the bubblers were active in collecting HTO from the exposure chamber is shown on the time scale. They were integrated measurements over the intervals shown. The model fits best at extended times where the intercepts with the "average-value" line segments are at the correct times. Additional uptake and release channels for short times, beyond those modeled, may be responsible for the early time disparity. A time lag of about 3 hours initially for, say, mixing would make the calculation agree very well with the experiment.

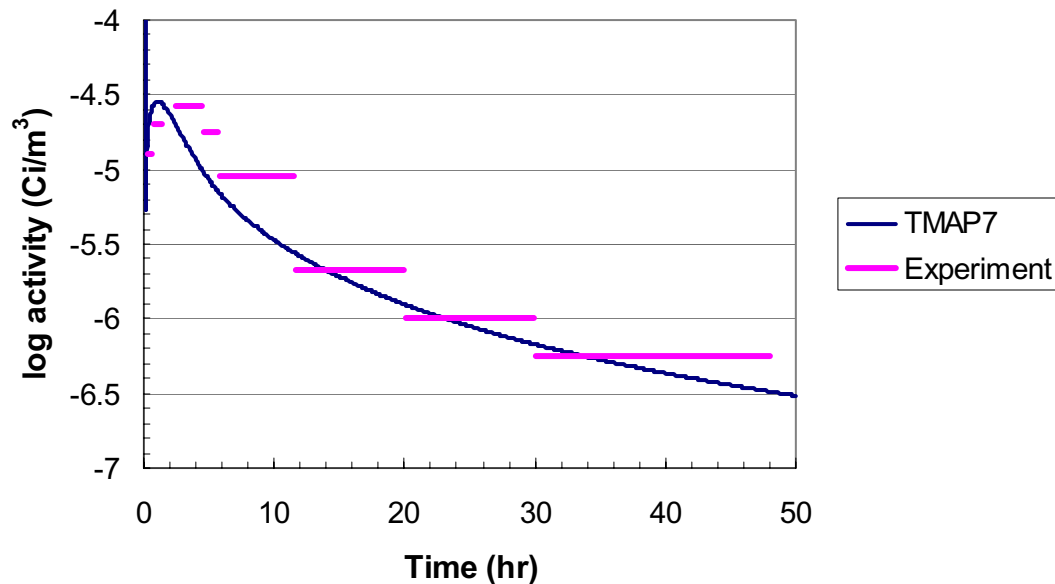


Figure 23. HTO Concentration in TSTA Exposure Chamber (Val-2c).

3.4 Problem 2d. Thermal Desorption Spectroscopy on Tungsten (Val-2d)

To exercise surface-law dependent diffusion boundary conditions and at the same time the multiple trapping capability, the experimental result of Hino et al.²³ was selected for approximation. In this experiment, H_3^+ was implanted at 5 keV and a flux of $1 \times 10^{19} \text{ H/m}^2\text{s}$ for

5,000 seconds into a polycrystalline tungsten foil 50 x 50 mm² and 0.1 mm thick at room temperature. Background pressure in the implantation chamber was 10⁻³ Pa while the implantation was going on and 10⁻⁵ Pa at other times. Following the implantation, the sample was subjected to thermal desorption spectroscopy by heating under vacuum at 50 K/min to 1,273 K and then held at that temperature for several minutes.

We modeled this system with TMAP7 using the structure of Figure 18. We first supposed that the evacuation pump was rated at 50 L/min or 8.33 x 10⁻⁴ m³/s. We then estimated the implantation beam area from the pumping rate, the indicated exposure chamber pressure during implantation of 10⁻³ Pa of H₂, and the implantation flux equivalent of 5 x 10¹⁸ H₂/m²/s, which is effectively all re-emitted from the target during irradiation.

$$A = \frac{PQ}{\phi kT} = \frac{\left(1 \times 10^{-3} \frac{J}{m^3}\right) \left(8.33 \times 10^{-4} \frac{m^3}{s}\right)}{\left(5 \times 10^{18} \frac{H_2}{m^2 s}\right) \left(1.3803 \times 10^{-23} \frac{J}{K}\right) (300 K)} = (4.025 \times 10^{-5} m^2) \quad (64)$$

We adjusted the enclosure volume to approximate the time constant of 625 seconds evident in the decay of what is effectively a pressure measurement at the end of the experiment. The volume chosen was 0.12 m³, which seems reasonable from the sketch provided in the report.²³

The test chamber was defined for this problem as a *functional* enclosure. We supposed that at least four pumping time constants (576 s) elapsed after the implantation ended before the thermal ramp began. We assumed the chamber would have a preprogrammed temperature of 300 K for 7,500 seconds followed by a ramp to 1,273 K at a ramp rate of 50 K/min. The vacuum pump is represented by a *boundary* enclosure (Encl 2) held at 10⁻⁸ Pa. Gas leakage from the ion source and elsewhere was modeled as a 10⁻⁵ Pa boundary enclosure (Encl 3) with flow to the exposure chamber at the vacuum pumping rate. The resulting pressure of background gases in the implantation chamber is thus realistic and reflects the 625-s settling time at the end of the experiment.

On the basis of TRIM²⁴²⁵ calculations, implantation was assumed to follow a normal distribution, peaking at 4.6 nm below the surface and having a scatter or characteristic half width of 3 nm. Implantation was active for 5000 seconds and then terminated.

The diffusion boundary condition employed was the *surfdep* or surface law dependent with the following parameter values

atomic hydrogen, H

ν = Debye frequency, 8.4E-12 (s⁻¹)

E_c = surface binding energy, -0.8 (eV)

E_s = solution enthalpy, 1.04 (eV)

P_c = combination probability, 1.0 (to form H₂)

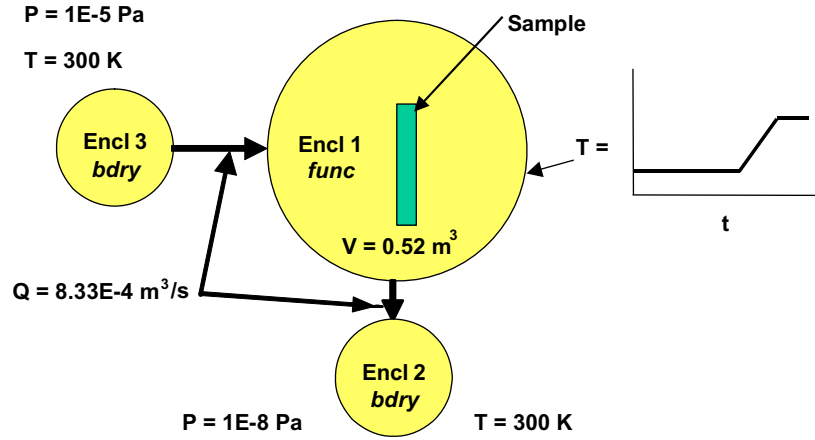


Figure 24. Schematic of system used to model experiments of Hino et al.²³

surface hydrogen, H₂

ν_o = Debye frequency, 8.4E-12 (s⁻¹)

E_c = surface binding energy, -0.1 (eV)

E_x = surface barrier energy, 0.05 (eV)

E_b = dissociation energy, 0.05 (eV)

M_m = molecular mass, 2.0 (amu)

P_c = formation probability, 1.0 (when H finds H)

ν_s = stoichiometric exponent, 0.5 (molecules per H combined)

For solubility of H in W, we use the value given by Frauenfelder.²⁶

$$S = 1.83 \times 10^{24} \left(\frac{H}{m^3} \right) \exp \left(- \frac{1.04 \text{ eV}}{RT} \right) \quad (65)$$

Diffusivity used for H through W was the normally accepted Frauenfelder value.²⁶

$$D = 4.1 \times 10^{-7} \left(\frac{m^2}{s} \right) \exp \left(- \frac{0.39 \text{ eV}}{RT} \right) \quad (66)$$

H₂ was considered insoluble in W and therefore had no diffusivity through the bulk. However, the surface diffusivity was taken to be

$$D = 4.1 \times 10^{-7} \left(\frac{m^2}{s} \right) \exp \left(- \frac{0.1 \text{ eV}}{RT} \right) \quad (67)$$

Three traps were assumed in the sample. Trap concentrations and distributions were considered adjustable parameters while energies were determined by TDS peak temperatures. The first was assumed to be associated with implantation (damage and precipitation) and to be normally distributed with a peak at 4.6 nm and a characteristic width of 10 nm, consistent with the observations of Haasz et al.²⁷ that damage zone exceeds the implantation depth. Its trap energy was adjusted, based on the temperature of the first peak, to be 1.0 eV, and it was assumed to be 7% atom fraction and uniform throughout the implantation region. A slightly lower value, say 6% or 5.5% would have given a better fit with the measured data. The second was a uniform trap, probably associated with dislocations and was assigned a trap release energy of 1.35 eV, typical of but slightly higher than that seen by Anderl et al.²⁸ Its concentration was adjusted to 1.38E-05 atom fraction. The third trap was also assumed to be uniformly distributed and to have a trapping energy of 2.7 eV, nearly the same as the deep trap seen by Frauenfelder²⁶ with a concentration of 5.9E-06 atom fraction. These were assumed to be approximately 90% filled at the start of the experiment, the hydrogen having come from air water vapor following previous anneals. Assuming emptier traps results in somewhat lower emissions, especially for the intermediate energy trap, because they do not completely fill in the 5,000 seconds of implantation. Higher diffusivity would correct that.

Even though the measured flux in the experiment was given in terms of a molecular flux density from the surface of the sample, it is likely that this corresponded to a pressure in the vicinity of the sample. Hence, the results reported here are those calculated for the enclosure H₂ pressure. The implantation flux of 10¹⁹ H/m²/s generated a molecular gas (H₂) pressure of 1 x 10⁻³ Pa during implantation. Therefore, a peak emission rate of 10¹⁸ H₂/m²/s during thermal desorption would correspond to a molecular gas pressure of 1 x 10⁻⁴ Pa. The computed results are shown together with the scaled Hino data in Figure 19.

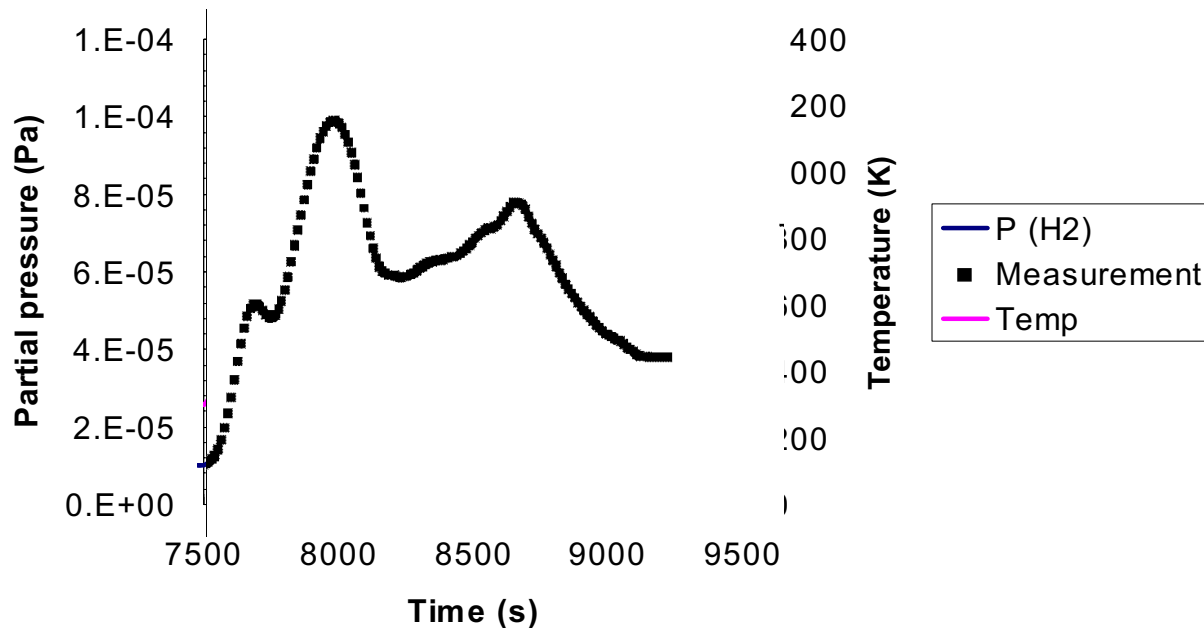


Figure 25. Comparison of calculated with experimental results for Hino's experiment with implantation and thermal desorption of tungsten (Val-2d).

The fit with the Hino et al. data is not exact because of several factors, the most prominent of which is probably that there are more than three traps. It appears that there may be several between 1.35 and 2.7 eV.

3.5 Problem 2e. Co-permeation of H₂ and D₂ through Pd (Val-2e)

This problem was selected to demonstrate a non-classical solution law boundary condition with molecular exchange as well as combined solution-law and recombination limited boundary conditions. It comes from work reported by Kizu et al.²⁹ on experiments in which H₂ and D₂ were allowed to permeate through thin Pd membranes either separately or together. The tests resulted in the formation of HD, both on the upstream side and on the downstream side of the membrane.

The experimental apparatus consisted of two vacuum chambers separated by a Pd membrane which was $1.8 \times 10^{-4} \text{ m}^2$ in area and either 0.025 mm or 0.05 mm thick, depending on the test. The membrane was clamped on each side by a copper gasket, and it may reasonably be inferred that the only means of transfer of gas from one chamber to the other was by diffusion through the membrane. Temperatures in the membrane were controlled between 820 and 870 K by means of an electric resistance heater surrounding the membrane and a thermocouple touching the membrane. Gas was introduced into one of the chambers from regulated supply bottles at various compositions and pressures. Here, we refer to that chamber as the upstream chamber. The base pressure on both upstream and downstream chambers was maintained at less than 10^{-6} Pa by a combination of turbomolecular pump and rotary backing pump on each side. Pressure was indicated by an ion gage on each side, and gas composition was measured with a quadrupole mass spectrometer. Flow rates through the membrane were determined by pressure increases in the downstream chamber at fixed pumping rate of $0.1 \text{ m}^3/\text{s}$.

The first tests reported were permeation tests of D₂ alone through membranes of each thickness. For the thinner membrane, tests were conducted at both 825 K and 865 K whereas the 0.05-mm membrane was tested only at 825 K. These were performed to calibrate the permeability of the membranes to hydrogen isotopes. Figure 20 shows their experimental data for permeation flux, $J(\text{D}_2)$, as a function of upstream D₂ pressure, $P(\text{D}_2)$.

Also shown in Figure 26 are three “fit” lines. Kizu et al. observed that at low pressures the permeation flux is directly proportional to the upstream gas pressure. As pressure increases, the permeation flux falls off from that linear relationship and approaches a square root relationship. Here, the fit to the 0.05-mm data (825 K) is made using least-squares methods across the range of pressures measured, not just at the lower pressures where greater linearity is observed. The fit line to the 0.025-mm data (825 K) is not really a fit at all. It is simply the line from the 0.05-mm data multiplied by a factor of 2. It fits the data amazingly well, indicating that permeation through the membrane is diffusion-limited, not surface-limited. The fit line for the 865-K data, also an extrapolation, has the same slope (0.8958) as the previous two fit lines, but it is offset by a factor of 1.55 from the 0.05-mm (825 K) line. It does not fit the higher-pressure data as well as it does the low-pressure data, but it does suggest a permeability activation energy of 0.674 eV (7,818 K). The resulting equation for D₂ permeability in Pd is thus

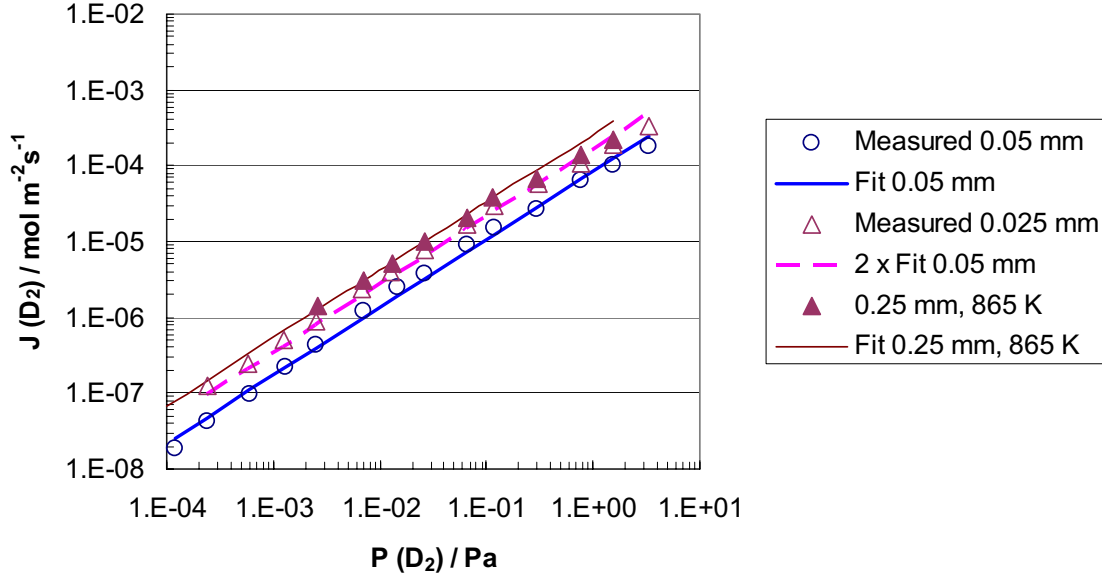


Figure 26. Permeability data of Kizu et al. for D₂ in Pd.

$$J = \frac{1.096 \times 10^{-4}}{L} P^{0.8958} \exp\left(-\frac{7818}{T}\right) \left(\frac{\text{mole}}{\text{m}^2 \text{s}}\right) \quad (68)$$

where

L = membrane thickness (m)

P = upstream pressure (Pa)

T = Temperature (K)

For the diffusion-limited regime, permeability is the product of solubility, S , and diffusivity, D , such that, approximately

$$J = \frac{C_0}{L} D = \frac{S P^\nu}{L} D = \frac{S_0 P^\nu D_0}{L} \exp\left[-\frac{(E_d + E_s)}{kT}\right] \quad (69)$$

where E_d and E_s are the diffusion activation energy and solution enthalpy, respectively. Comparing Eqs. (67) and (68), we see that

$$\nu = 0.8958$$

$$S_0 D_0 = 1.096 \times 10^{-4}$$

$$E_d + E_s = 7,818 \text{ k}$$

We can separate diffusivity and solubility by making use of the diffusivity of hydrogen in Pd given by Katz and Gulbransen³⁰ divided by $\sqrt{2}$ to account for isotopic effect on diffusivity

$$D_D = 3.048 \times 10^{-7} \exp\left(-\frac{2818}{T}\right) \left(\frac{\text{m}^2}{\text{s}}\right) \quad (70)$$

That leaves for deuterium

$$S_D = 179.6 \exp\left(-\frac{5000}{T}\right) \left(\frac{\text{mole}}{\text{m}^3 \text{Pa}^v}\right) = 1.082 \times 10^{26} \exp\left(-\frac{5000}{T}\right) \left(\frac{\text{atom}}{\text{m}^3 \text{Pa}^v}\right) \quad (71)$$

Kizu et al.²⁹ note that the solubility of hydrogen in Pd is about 1.23 times as great as the solubility of deuterium in Pd at the temperature of their experiments.

Next, we construct a model for TMAP7 simulation of this experiment. We consider two functional enclosures, each with an estimated volume of 0.1 m^3 , separated by a diffusion segment of thickness L and area $1.8 \times 10^{-4} \text{ m}^2$. This is illustrated in Figure 27.

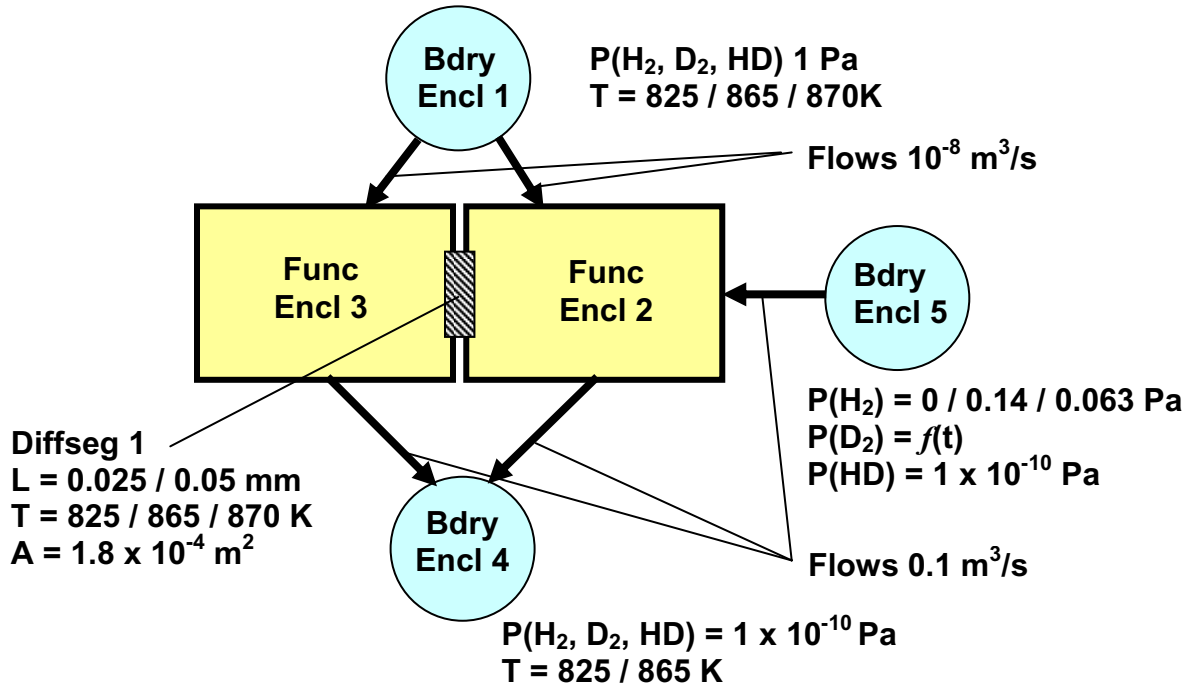


Figure 27. TMAP7 model of experimental system of Kizu et al.

Boundary enclosure 1 is the source of background pressure to the experimental system. Boundary enclosure 4 is the vacuum pumping system that provides a sink for all system flows. Boundary enclosure 5 is the gas feed to the upstream experimental chamber, functional enclosure 2. Depending on the experiment, the feed pressure of H_2 is 0, 0.14 Pa, or 0.063 Pa. Combined with the evacuation to boundary enclosure 4, this provides the upstream H_2 pressure for permeation. The D_2 pressure is a stepped function of time, one step corresponding to each of the data points in the data plots of Kizu et al. Steps are arbitrarily set at 100 s, but equilibrium is achieved in times much shorter than that. No HD is fed into the upstream experimental chamber, in keeping with the experimental setup given by Kizu et al. For the solution-law (*lawdep*) boundary conditions, it is assumed that HD is formed in accordance with the laws of chemical equilibrium.

$$P_{HD} = 2\sqrt{P_{H_2} P_{D_2}} \quad (72)$$

Under these conditions, one specifies the homonuclear gas pressures, and TMAP7 determines what the corresponding heteronuclear gas pressure must be. Thus, for this analysis, one specifies effective deuterium pressure P_D , where

$$P_D = P_{D_2} + \frac{P_{HD}}{2} \quad (73)$$

and similarly for P_H .

In the experiments, even though H_2 gas flow is fixed, it is not P_{H_2} that is fixed but P_H , because as the D_2 pressure increases, some of the H_2 will be converted to HD. Under equilibrium (*lawdep*) conditions, it may be shown that

$$P_{HD} = \frac{2P_H P_D}{P_H + P_D} \quad (74)$$

which, with Eq. (73) leads to

$$P_{D_2} = \frac{P_D^2}{P_H + P_D} \quad (75)$$

Likewise, for the calculation, we will not have constant P_{H_2} , but P_{H_2} will vary according to

$$P_{H_2} = \frac{P_H^2}{P_H + P_D} \quad (76)$$

For the non-equilibrium (*ratedep*) boundary condition, the pressure in enclosure 5 will be the specified P_D , and the code will determine the enclosure pressures of H_2 , D_2 , and HD from the pumping, dissociation, and recombination rates.

For the downstream chamber, functional enclosure 3 under *lawdep* conditions, HD is formed together with H_2 and D_2 in chemical equilibrium from diffusing H and D. Under *ratedep* conditions, again the relative dissociation and recombination rates together with the convective flows determine the partial pressures in enclosure 3.

We first replicate the calibration experiments shown in Figure 26 using input files [Val-2ea.inp](#), [Val-2eb.inp](#), and [Val-2ec.inp](#) for the three cases shown in Figure 26. Results are in Figure 28. The results are almost as good as the approximations for the permeability in Figure 26.

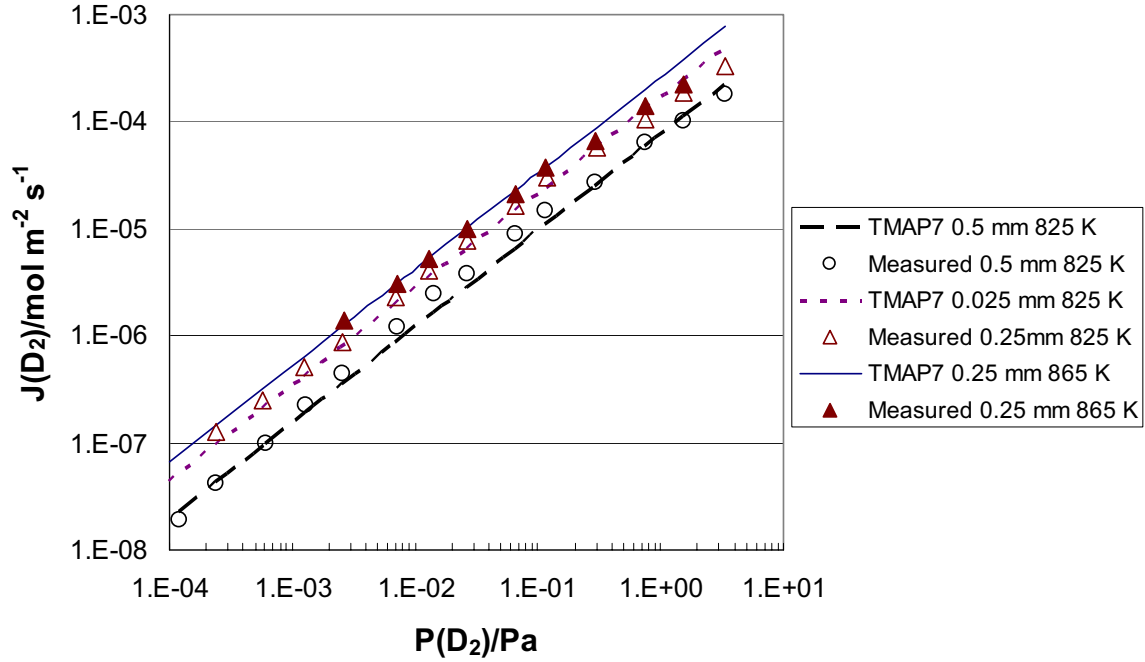


Figure 28. Comparison of TMAP7 permeation calculations with permeation data of Kizu et al. for D_2 only under *lawdep* boundary conditions using the solubility of Eq. (71) (Val-2ea, Val-2eb, Val-2ec).

In modeling the co-permeation of H and D, we first apply the *lawdep* boundary condition in which we apply H_2 through enclosure 5 at pressures indicated by Eq. (76) for pre-selected values of P_H (0.063 Pa) and P_D corresponding to the abscissa values of the data in Kizu et al.²⁹ D_2 is also added at pressures given by Eq. (75) for the same P_H and set of P_D values for the experiment on a 0.025-mm membrane at 870 K (Val-2ed.inp). The results of that computation are compared with the experimental data in Figure 29.

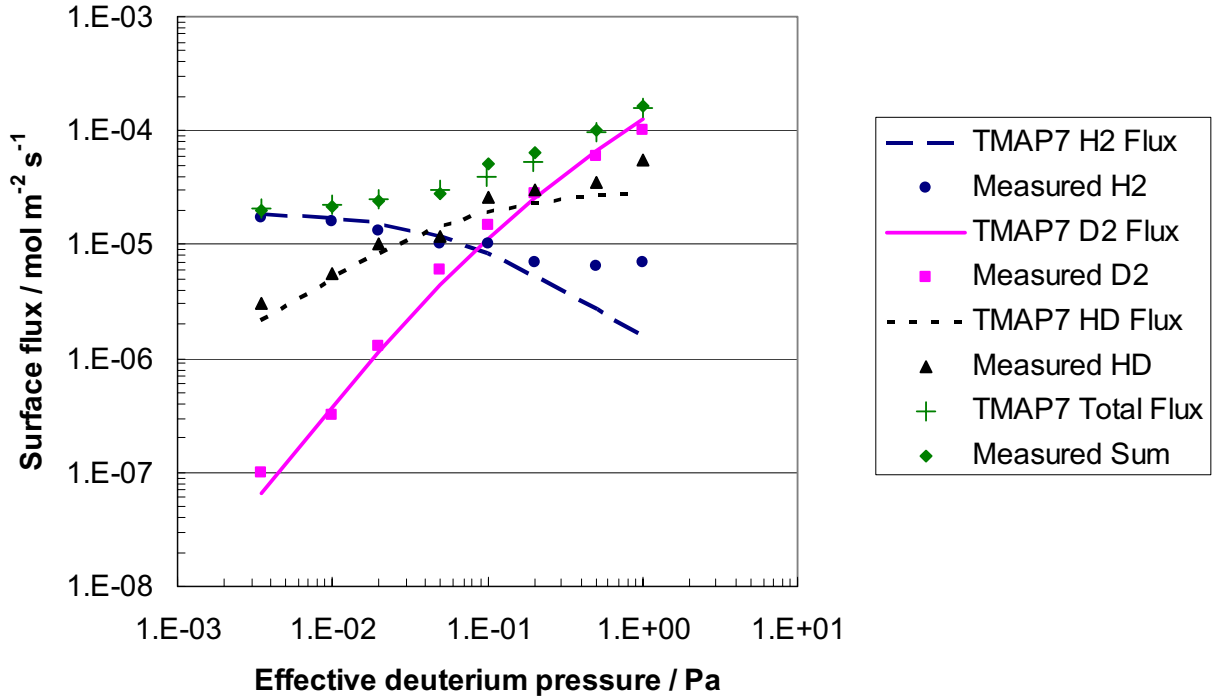


Figure 29. Comparison of TMAP7 results using a lawdep boundary condition on each side of the membrane with the experiments of Kizu et al. (Val-2ed).

It is evident that while the H₂ permeation calculated at low effective deuterium pressures agree well with the experimental data, they do not agree at higher deuterium pressures. Likewise, HD release rates are under-predicted at high D₂ pressures. D₂ release rates agree reasonably well at higher and lower pressures but slightly less well at intermediate pressures.

For additional perspective, we next changed the diffusion boundary condition to the *ratedep* mode in which dissociation and recombination take place independently ([Val-2ee.inp](#)). We use for the dissociation rate coefficient the molecular arrival rate at the surface

$$K_d = \frac{1}{\sqrt{2\pi M k T}} = \frac{2.6276 \times 10^{24}}{\sqrt{M T}} \left(\frac{\text{molecule}}{m^2 Pa} \right) \quad (77)$$

where M is the species molecular weight in amu. For the recombination coefficient, we use the relationship from Sieverts' law that

$$K_r = \frac{K_d}{S^2} \quad (78)$$

For deuterium, using S from Eq. (71),

$$K_{r_{D_2}} = \frac{1.994 \times 10^{-28}}{\sqrt{M T}} \exp\left(\frac{10000}{T}\right) \left(\frac{m^4}{s} \right) \quad (79)$$

while for hydrogen with its higher solubility

$$K_{r_{H_2}} = \frac{1.318 \times 10^{-28}}{\sqrt{MT}} \exp\left(\frac{10000}{T}\right) \left(\frac{m^4}{s}\right) \quad (80)$$

For HD we use the average of these two. The results from that computation are as shown in Figure 30.

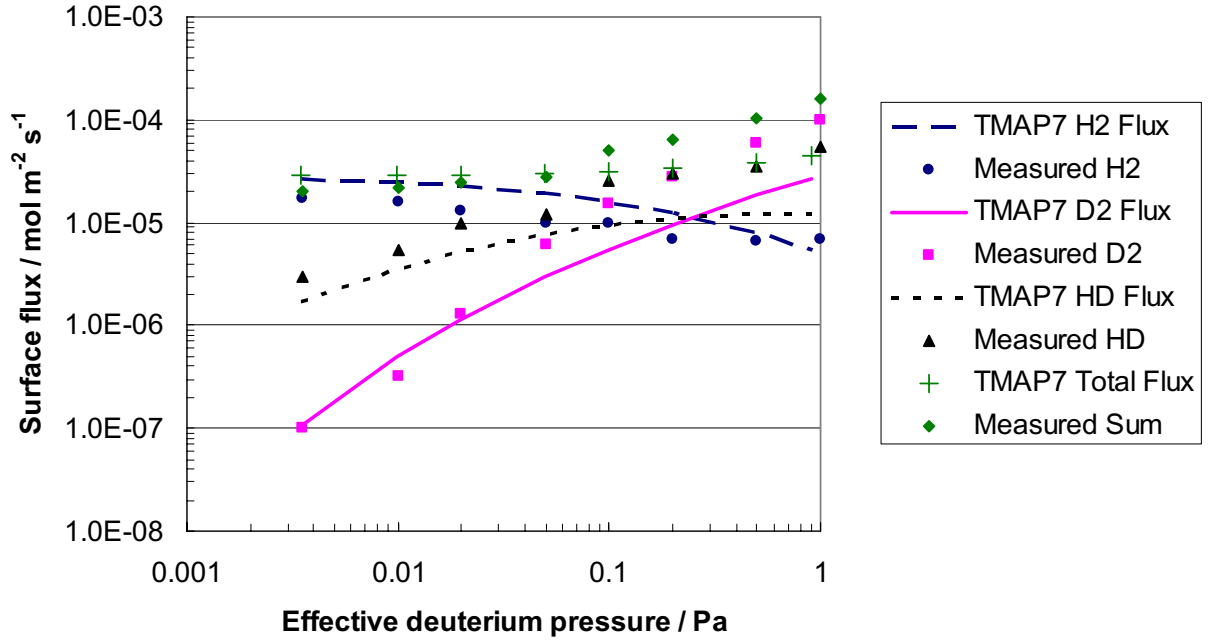


Figure 30. Comparison of TMAP7 calculation with simple *ratedep* boundary conditions with the values measured by Kizu et al. (Val-2eea).

Here, H₂ permeation at low D₂ pressures is over-predicted but it improves at higher deuterium pressures. Agreement for HD is better at low pressures than high. D₂ permeation is good at low D₂ pressures but it under-predicts markedly at high pressures. Total permeation rate is good at intermediate pressures but it is over-predicted at low pressures and under-predicted at high pressures. It appears that more deuterium and hydrogen are getting into the upstream face of the membrane at higher pressures than are predicted by the model. Increasing K_d to the full arrival rate has little effect on results. Reducing K_d to 0.1 of the molecular arrival rate and using the same approach shifts the permeation curves upward, as shown in Figure 31.

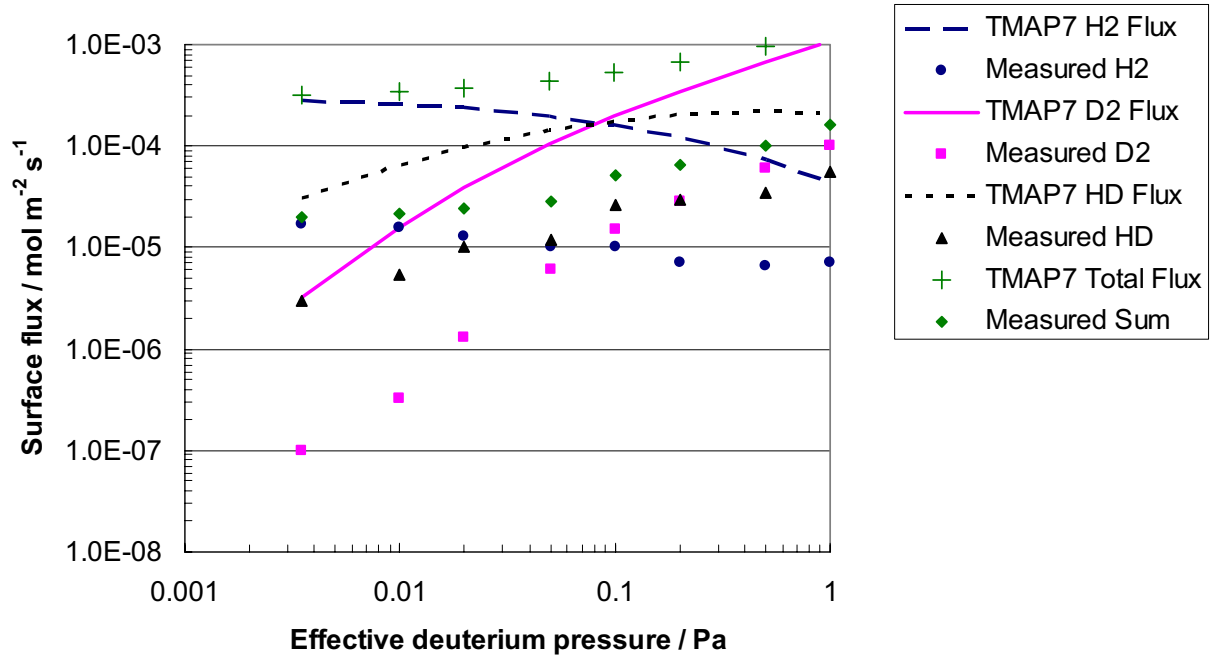


Figure 31. TMAP7 results from a 5-fold reduction in K_d and K_r (Val-2eeb).

Though the fit is not as good as the *lawdep* solution, these results are consistent with the observations of Kizu et al. that permeation appears to be nearly first-order in P at low pressures but tends to become proportional to $P^{1/2}$ as driving pressure increases. For a variation, the problem was rerun with a *lawdep* upstream diffusion boundary condition and a *ratedep* boundary condition downstream (Val-2ef). The results are shown in Figure 32. The fit is not particularly good anywhere.

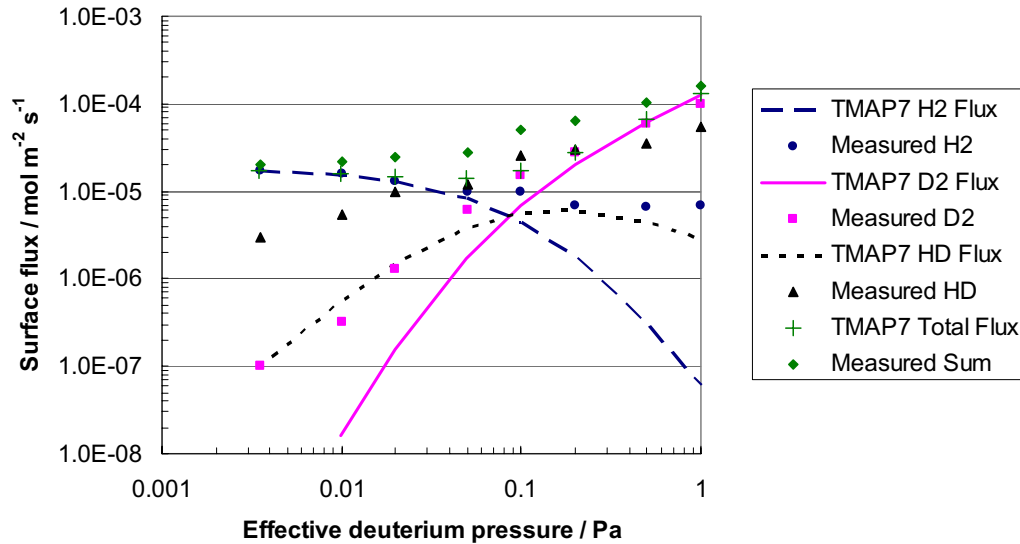


Figure 32. Comparison of TMAP7 calculation for *lawdep* boundary condition upstream and *ratedep* boundary condition downstream with measurements made by Kizu et al. (Val-2ef).

4.0 CONCLUSIONS

In the course of the work performed here, the TMAP7 code has been demonstrated in a wide variety of applications. Many of these are contrived problems for which analytical solutions are available. Agreement between solutions calculated by TMAP7 and those generated in a Microsoft Excel™ spreadsheet is excellent. A second group of problems constitute replications of actual experiments, the results of which appear in published journals. By making use of accepted values of transport parameters and some fitting constant values, it has been shown that TMAP7 gives results in good agreement with actual measurements. These two groups of exercises constitute the verification and validation of the TMAP7 code.

The major challenge in assembling the computational models is finding the necessary parameters for the various property values needed in the code. A further challenge with TMAP7 is one faced by many such codes, numerical convergence. This is managed with various control parameters to adjust the damping in convergence iteration.

TMAP7 represents a significant step forward in modeling gas interaction with structures and in enclosures.

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APPENDIX A SPECIES EQUILIBRATION MODEL

Suppose that two homonuclear diatomic molecular species, A_2 and B_2 , are in a volume V , and at time $t = 0$, are allowed to contact a catalytic surface of area S that supports the reaction



Assume further that the molecular species have the same mass and chemical properties such that there is no enthalpy change associated with this reaction and only configurational entropy is driving the reaction. Then

$$\Delta G_f = -T\Delta s_f = -RT \ln 2 \quad (A-2)$$

The equilibrium constant for reaction (A-1) is then

$$K_{eq} = \exp\left(-\frac{\Delta G_f}{RT}\right) = 2 \quad (A-3)$$

The law of mass action then requires that in equilibrium,

$$\frac{[AB]}{[A_2]^{\frac{1}{2}}[B_2]^{\frac{1}{2}}} = 2 \quad (A-4)$$

or equivalently

$$P_{AB} = 2 P_{A_2}^{\frac{1}{2}} P_{B_2}^{\frac{1}{2}} \quad (A-5)$$

The AB molecules come from the dissociation of A_2 and B_2 molecules such that for starting pressures $P_{A_2}^0$ and $P_{B_2}^0$, it must also be true that at equilibrium

$$P_{AB} = 2 \frac{P_{A_2}^0 P_{B_2}^0}{P_{A_2}^0 + P_{B_2}^0} \quad (A-6)$$

Two different approaches to the dynamics of the equilibration process will now be explored, one corresponding to *ratedep* boundary conditions and the other to *surfdep* conditions.

***Ratedep* Conditions**

At equilibrium, when Sieverts' law applies, for atom concentrations C_A and C_B at the surface,

$$\begin{aligned} C_A &= K_s \sqrt{P_{A_2}} \\ C_B &= K_s \sqrt{P_{B_2}} \end{aligned} \quad (A-7)$$

where K_s is the Sieverts' solubility. Because of the assumed equality of chemistry, K_s will be the same for each homonuclear species. We expect also that under equilibrium conditions

$$K_d P_{A_2} = K_r C_A^2 \quad (\text{A-8})$$

where K_d is the dissociation coefficient and K_r is the recombination coefficient. That leads to

$$K_d = K_s^2 K_r \quad (\text{A-9})$$

We expect further for the heteronuclear species

$$K_d P_{AB} = K_{rAB} C_A C_B \quad (\text{A-10})$$

Under *ratedep* conditions, equilibrium is not assumed, but the relationships between the coefficients are maintained. Under these assumed conditions, the dissociation coefficients for both AB and A_2 or B_2 molecules should be identical. However, because two different microscopic processes can produce AB (A jumping to find B and B jumping to find A) and only one (A finding A) can form A_2 , and similarly for B_2 , we expect K_{rAB} to be twice K_r for the homonuclear molecules.

We first write conservation equations for the surface species, C_A and C_B .

$$\begin{aligned} C_A(C_A + C_B)2K_r &= K_d(2P_{A_2} + P_{AB}) \\ C_B(C_A + C_B)2K_r &= K_d(2P_{B_2} + P_{AB}) \end{aligned} \quad (\text{A-11})$$

Adding these together and applying the conservation of gas atoms in the enclosure gives

$$(C_A + C_B)^2 = K_s^2 (P_{A_2}^0 + P_{B_2}^0) \quad (\text{A-12})$$

This requires that C_A and C_B are both constant.

The current of AB molecules from surface S from volume V is the rate of change of those molecules in the enclosure.

$$\frac{dN_{AB}}{dt} = S(2K_r C_A C_B - K_d P_{AB}) \quad (\text{A-13})$$

Here, N_{AB} is the number of molecules of species AB in the enclosure. Solving Equations (A-11) for $C_A C_B$, we find that

$$C_A C_B = \frac{K_d P_{A_2}^0 P_{B_2}^0}{2K_r (P_{A_2}^0 + P_{B_2}^0)} \quad (\text{A-14})$$

Then, Equation (A-13) becomes

$$\frac{dP_{AB}}{dt} = \frac{SK_d kT}{V} \left(\frac{2P_{A_2}^0 P_{B_2}^0}{(P_{A_2}^0 + P_{B_2}^0)} - P_{AB} \right) \quad (\text{A-15})$$

Equation (A-15) is solved by

$$P_{AB} = \frac{2P_{A_2}^0 P_{B_2}^0}{(P_{A_2}^0 + P_{B_2}^0)} \left[1 - \exp\left(-\frac{SK_d kT}{V} t\right) \right] \quad (\text{A-12})$$

Surfdep Conditions

When *surfdep* conditions apply, there are no assumptions about equilibrium except in the steady state. Then, the surface concentration of molecules is directly proportional to the gas over-pressure and we define a deposition rate constant by.

$$\hat{K}_d = \frac{1}{\sqrt{2\pi MkT}} \exp\left(-\frac{E_x}{kT}\right) \quad (\text{A-14})$$

where M is the mass of any of the species molecules, assuming all are equal, and E_x is the adsorption barrier energy. For release of the molecular species from the surface,

$$\hat{K}_r = \frac{\nu_o}{6} \exp\left(\frac{E_c - E_x}{kT}\right) \quad (\text{A-15})$$

Here, ν_o is the Debye frequency, E_c is the surface binding energy, and the factor of 6 accounts for the probability that a given phonon will be directed away from the surface. At steady-state, the flux to the surface will be balanced by flux from the surface, and surface concentration will be related to the gas over-pressure by

$$C_{m_s} = P_m \frac{\hat{K}_d}{\hat{K}_r} = \frac{6P_m}{\nu_o \sqrt{2\pi MkT}} \exp\left(-\frac{E_c}{kT}\right) \quad (\text{A-16})$$

The conversion of A_2 and B_2 molecules to AB molecules requires several steps. First, homonuclear molecules in the gas must get to the surface. Next, they must dissociate. Then the individual surface atoms must migrate to sites where they encounter their conjugates. Here we assume there is a probability of unity of their combination once they find each other. Finally, the AB molecule must leave the surface and return to the gas. We write equations for species continuity at the surface.

$$C_{AB}(\hat{K}_r + \hat{K}_b) = P_{AB} \hat{K}_d + C_A C_B (2D_s \lambda) \quad (\text{A-17})$$

$$C_{A_2}(\hat{K}_r + \hat{K}_b) = P_{A_2} \hat{K}_d + C_A^2 (D_s \lambda) \quad (\text{A-18})$$

$$C_{B_2}(\hat{K}_r + \hat{K}_b) = P_{B_2} \hat{K}_d + C_B^2 (D_s \lambda) \quad (\text{A-19})$$

$$C_A [(C_A + C_B) 2D_s \lambda] = (C_{AB} + 2C_{A_2}) K_b \quad (\text{A-20})$$

$$C_B [(C_A + C_B) 2D_s \lambda] = (C_{AB} + 2C_{B_2}) K_b \quad (\text{A-21})$$

In these equations, the dissociation rate for molecules at the surface is given by

$$K_b = \exp\left(-\frac{E_b}{kT}\right) \quad (\text{A-22})$$

where E_b is the dissociation activation energy, D_s is the surface diffusivity of the atomic species, and λ is the lattice constant, assumed to be the reciprocal cube root of the lattice density. K_b is assumed equal for all molecular species, and D_s is assumed to be the same for all atomic species.

We may combine Equations (A-17) to (A-21) to find that

$$P_{total} = P_{A_2} + P_{B_2} + P_{AB} = (C_A + C_B)^2 \left(\frac{\hat{K}_r}{K_b} \right) \frac{D\lambda}{\hat{K}_d} \quad (A-23)$$

This is reminiscent of Sieverts' law. With the conservation law for atoms in the gas

$$P_{AB} = 2(P_{A_2}^0 - P_{A_2}) = 2(P_{B_2}^0 - P_{B_2}) \quad (A-24)$$

Equation (A-23) becomes

$$(C_A + C_B)^2 = (P_{A_2}^0 + P_{B_2}^0) \frac{\hat{K}_d}{D_s \lambda} \frac{K_b}{\hat{K}_r} \quad (A-25)$$

Note that no assumption has been made regarding steady state. Because the sum of the concentrations C_A and C_B is constant in time for this problem, either the individual concentrations must both be constant or a change in one must be the negative of a change in the other. The latter case is not consistent with the definition of present problem. Therefore, they must both be constant. Then, from statistical considerations, the molecular formation rates must be the same as they are in steady state.

The process that converts dissociation products to AB molecules is the recombination step while the net destruction rate is dissociation. Hence

$$\frac{dN_{AB}}{dt} = S(C_A C_B 2D_s \lambda - C_{AB} K_b) \quad (A-26)$$

Equation (A-17) must hold at all times such that if we solve it for C_{AB} and substitute the result into Equation (A-26) we get, successively

$$\begin{aligned} \frac{dN_{AB}}{dt} &= S \left(C_A C_B 2D_s \lambda - K_b \frac{P_{AB} \hat{K}_d + C_A C_B 2D_s \lambda}{\hat{K}_r + K_b} \right) \\ \frac{dP_{AB}}{dt} &= \frac{SkT}{V} \left[C_A C_B 2D_s \lambda \left(1 - \frac{K_b}{\hat{K}_r + K_b} \right) - P_{AB} \frac{\hat{K}_d K_b}{\hat{K}_r + K_b} \right] \\ \frac{dP_{AB}}{dt} &= \frac{SkT}{V} \frac{\hat{K}_d K_b}{\hat{K}_r + K_b} \left[C_A C_B 2D\lambda \left(\frac{\hat{K}_r}{\hat{K}_d K_b} \right) - P_{AB} \right] \end{aligned} \quad (A-27)$$

This is solved by the expression

$$P_{AB} = C_A C_B 2D\lambda \frac{\hat{K}_r}{\hat{K}_d K_b} \left[1 - \exp \left(- \frac{SkT t}{V} \frac{\hat{K}_d K_b}{\hat{K}_r + K_b} \right) \right] \quad (A-28)$$

It may be shown, again using Equations (A-17) to (A-21), that this is equivalent to

$$P_{AB} = 2 \frac{P_{A_2}^0 P_{B_2}^0}{P_{A_2}^0 + P_{B_2}^0} \left[1 - \exp \left(- \frac{t}{\tau} \right) \right] \quad (A-29)$$

where

$$\tau = \frac{V(\hat{K}_r + K_b)}{SkT \hat{K}_d K_b} \tag{A-30}$$

APPENDIX B

PROBLEM INPUT FILE LISTINGS

In this appendix are the input file listings used in the demonstration problems in Sections 2 and 3. These may be used as starting points for individual problems by the user.

Problem 1a: Diffusion from a Depleting Source (Val-1a, see p. 2)

```

title input
  Validation Problem #1a. Tritium diffusion through SiC layer
  with depleting source at 2100C. No solubility or trapping included.
end of title input
$ -----
main input
$ -----
dspcnme=t,end
espcnme=ts,end
segnds=9,end
nbrencl=2,end
end of main input
$ -----
enclosure input
$ -----
start func,1,end
etemp=2373.0,end
esppres=ts,1.0e6,end
evol=5.2e-11,end
$
start bdry,2
etemp=2373.0,end
esppres=ts,0.0,end
end of enclosure input
$ -----
thermal input
$ -----
start thermseg,end
delx=0.0,3.0e-6,6*5e-6,0.,end
tempd=9*2373.0,end          $ Initial temperatures=(K)
end of thermal input
$ -----
diffusion input
$ -----
start diffseg,end
$ Sample [DIFFSEG 1]
nbrden=4.832e28,end
concd=t,9*0.0,end
qstrdr=t,equ,3,end          $ Q*/R for Soret effect unknown
dcoef=t,equ,1,end          $ Diffusion coeff (m2/s)
srcsd=t,equ,3,srcpf,9*0.0,end
difbcl=lawdep,encl,1,dspc,t,ts,pexp,1.0,solcon,equ,2,end
difbcr=sconc,dspc,t,conc,const,0.0,end
surfa=2.16e-6,end
end of diffusion input
$ -----
equation input
$ -----
$ (1) Diffusion coefficient
y=1.58e-4*exp(-308000.0/(8.314*temp)),end
$ (2) Solubility constant
y=7.244e22/temp,end
$ (3) Soret coefficient
y=0.0,end
end of equation input

```



```

$ -----
table input
$ -----
end of table input
$ -----
control input
$ -----
time=0.0,end           $ initial time
tstep=0.01,end         $ time step = 10 msec
timend=140.001,end     $ the last time computed (s)
nprint=1000,end        $ print every 10 seconds
itermx=2000,end
delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
end of control input
$
plot input
nplot=100,end          $ makes plotfile entry every 1 sec
plotseg=1,end          $ segments for which plot info is needed
plotenc1=1,2,end       $ enclosures for which plot info is needed
dname=t,end            $ diffusing species for which plot info is needed
ename=ts,end           $ enclosure species for which plot info is needed
dplot=sflux,moblinv,end
eplot=press,end        $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1b: Diffusion in a Semi-Infinite Slab with Constant-Source Boundary (Val-1b, see p. 5)

```
title input
  Validation Problem #1a - 2100 C --No solubility or trapping.
  Tritium diffusion through semi-infinite layer w/ constant source
end of title input
$
main input
dspcnme=t,end
espcnme=ts,end
segnds=200,end
nbrencl=2,end
end of main input
$
enclosure input
start bdry,1,end
etemp=2373.0,end
esppres=ts,1.0e6,end
$
start bdry,2
etemp=2373.0,end
esppres=ts,0.0,end
end of enclosure input
$
thermal input
start thermseg,end
$delx=0.0,.001,.005,.01,.05,.1,.5,1.,5.,89*10.,0.0,end
delx = 0.0,198*0.1,0.0,end
tempd=200*2373.0,end
end of thermal input
$
diffusion input
start diffseg,end
nbrden=4.832e28,end
concd=t,200*0.0,end
qstrdr=t,equ,2,end
dcoef=t,const,1.0,end
srcsd=t,const,0.0,srcpf,200*0.0,end
difbcl=sconc,dspc,t,conc,const,1.0,end
difbcr=sconc,dspc,t,conc,const,0.0,end
surfa=1.0,end
$
end of diffusion input
$
equation input
end of equation input
$
table input
end of table input
$
control input
time=0.0,end
tstep=0.01,end
timend=50.0,end
nprint=100,end
itermx=20000,end
$ time step = 10 msec
$ after implantation and desorption
$ print every second
```

```

delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
end of control input
$
plot input
nplot=100,end          $ makes plotfile entry every 1 sec
plotseg=1,end          $ segments for which plot info is needed
plotenc1=1,2,end       $ enclosures for which plot info is needed
dname=t,end            $ diffusing species for which plot info is needed
ename=ts,end           $ enclosure species for which plot info is needed
dplot=sflux,end
eplot=end              $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1c: Diffusion in a Partially Preloaded Semi-Infinite Slab (Val-1c, see p. 8)

```
title input
  Validation Problem #3 - Transient Concentration for semi-infinite,
  partially preloaded slab with both boundaries at 0 Concentration
  T = 2100 K
end of title input
$ -----
main input
$ -----
dspcnme=td,end
espcnme=t,end
segnds=99,end
nbrencl=2,end
end of main input
$
$ -----
enclosure input
$ -----
start bdry,1,end
etemp=2373.0,end
esppres=t,0.0,end
$
start bdry,2,end
etemp=2373.0,end
esppres=t,0.0,end
$
end of enclosure input
$
$ -----
thermal input
$ -----
start thermseg,end
delx=0.0,10*1.0,0.5,5*1.0,5*5.0,76*10.0,0.0,end
tempd=99*2373.0,end
$
end of thermal input
$ -----
diffusion input
$ -----
start diffseg,end
$ Sample [DIFFSEG 1]
nbrden=4.832e28,end
concd=td,11*1.0,88*0.0,end
qstrdr=td,equ,2,end      $ Q*/R for Soret effect unknown
dcoef=td,equ,1,end      $ Diffusion coeff (m2/s)
srcsd=td,equ,2,srcpf,99*0.0,end
difbcl=sconc,dspc,td,conc,const,0.0,end
difbcr=sconc,dspc,td,conc,const,0.0,end
surfa=1.0,end
$
end of diffusion input
$
$ -----
equation input
$ -----
```

```

$ (1)
y=1.0,end
$ (2)
y=0.0,end
$
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
time=0.0,end
tstep=0.005,end          $ time step = 1 sec
timend=100.005,end
nprint=1000,end          $ print every 5 seconds
itermx=20000,end
delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
damp=1.0
end of control input
$
$ -----
plot input
$ -----
nplot=1000,end           $ makes plotfile entry every 5 sec
plotseg=1,end            $ segments for which plot info is needed
plotencl=1,end           $ enclosures for which plot info is needed
dname=td,end             $ diffusing species for which plot info is needed
ename=t,end              $ enclosure species for which plot info is needed
dplot=moblinv,sflux,sconc,end
eplot=diff,end           $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1da. Effective Diffusivity Trap (Val-1da, see p. 12)

```

title input
  Validation Problem #4a - Trapping in a slab of constant upstream
  concentration - trapping suspended
end of title input
$ -----
main input
$ -----
dspcnme=td,end
espcnme=t,end
segnds=22,end
nbrencl=2,end
end of main input
$
$ -----
enclosure input
$ -----
start bdry,1,end
etemp=1000.0,end
esppres=t,1.0,end
$
start bdry,2,end
etemp=1000.0,end
esppres=t,0.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
start thermseg,end
delx=0.0,20*0.05,0.0,end
tempd=22*1000.0,end
$
end of thermal input
$ -----
diffusion input
$ -----
start diffseg,end
$ Sample [DIFFSEG 1]
nbrden=3.162e22,end
concd=td,22*0.0,end
qstrdr=td,equ,2,end      $ Q*/R for Soret effect unknown
dcoef=td,equ,1,end      $ Diffusion coeff (m2/s)
srcsd=td,equ,2,srcpf,22*0.0,end
$trapping=ttyp,1,tconc,const,.1,tspc,td,alpht
$      equ,3,alphr,equ,4,ctrp,const,0.0,end
difbcl=sconc,dspc,td,conc,const,5.721e18,end
difbcr=sconc,dspc,td,conc,const,0.0,end
surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
$ (1)

```

```

y=1.0,end
$ (2)
y=0.0,end
$ (3)
y=1.0e15,end
$ (4)
y=1.0e13*exp(-100./temp),end
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
time=0.0,end
tstep=0.001,end
timend=3.0,end      $ after implantation and desorption
nprint=500,end      $ print every 0.5 seconds
itermx=200,end
delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
nplot=50 ,end      $ makes plotfile entry every 0.05 sec
plotseg=1,end      $ segments for which plot info is needed
plotencl=end       $ enclosures for which plot info is needed
dname=td,end       $ diffusing species for which plot info is needed
ename=end          $ enclosure species not needed
dplot=sflux,end
eplot=end          $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1db. Strong Trap (Val-1db, see p. 13)

```
title input
  Validation Problem #4b - Trapping in a slab of constant upstream
  concentration - strong-trapping limit
end of title input
$ -----
main input
$ -----
dspcnme=td,end
espcnme=t,end
segnds=22,end
nbrencl=2,end
end of main input
$
$ -----
enclosure input
$ -----
start bdry,1,end
etemp=1000.0,end
esppres=t,const,1.0,end
$
start bdry,2,end
etemp=1000.0,end
esppres=t,const,0.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
start thermseg,end
delx=0.0,20*0.05,0.0,end
tempd=22*1000.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
start diffseg,end
$ Sample [DIFFSEG 1]
nbrden=3.1622e22,end
concd=td,22*0.0,end
qstrdr=td,equ,2,end      $ Q*/R for Soret effect unknown
dcoef=td,equ,1,end      $ Diffusion coeff (m2/s)
srcsd=td,equ,2,srcpf,22*0.0,end
trapping=ttyp,1,tconc,const,.1,tspc,td,alph
    equ,3,alphr,equ,4,ctrp,const,0.0,end
difbcl=sconc,dspc,td,conc,const,3.1622e18,end
difbcr=sconc,dspc,td,conc,const,0.0,end
surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
$ (1)
```



```

y=1.0,end
$ (2)
y=0.0,end
$ (3)
y=1.0e15,end
$ (4)
y=1.0e13*exp(-15000./temp),end
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
time=0.0,end
tstep=0.1,end
timend=800.0,end
nprint=10,end
itermx=1000,end
delcmx=1.0e-6,end
bump=1.e-3,end
bound=8.0,end
damp=0.05,end
omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
nplot=1,end           $ makes plotfile entry every 1 sec
plotseg=1,end         $ segments for which plot info is needed
plotencl=end          $ enclosures for which plot info is needed
dname=td,end          $ diffusing species for which plot info is needed
ename=end             $ enclosure species for which plot info is needed
dplot=sflux,end
eplot=end             $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1dc. Multiple Trap (Val-1dc, see p. 14)

```
title input
  Validation Problem #4c - Trapping in a slab of constant upstream
  concentration - three distinct traps
end of title input
$ -----
main input
$ -----
dspcnme=td,end
espcnme=t,end
segnds=22,end
nbrencl=2,end
end of main input
$
$ -----
enclosure input
$ -----
start bdry,1,end
etemp=1000.0,end
esppres=t,const,1.0,end
$
start bdry,2,end
etemp=1000.0,end
esppres=t,const,0.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
start thermseg,end
delx=0.0,20*0.05,0.0,end
tempd=22*1000.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
start diffseg,end
$ Sample [DIFFSEG 1]
nbrden=3.1622e22,end
concd=td,22*0.0,end
qstrdr=td,equ,2,end      $ Q*/R for Soret effect unknown
dcoef=td,equ,1,end      $ Diffusion coeff (m2/s)
srcsd=td,equ,2,srcpf,22*0.0,end
trapping=ttyp,1,tconc,const,.1,tspc,td,alpht
      equ,3,alphr,equ,4,ctrp,const,0.0
      ttyp,2,tconc,const,.15,tspc,td,alpht
      equ,3,alphr,equ,5,ctrp,const,0.0
      ttyp,3,tconc,const,.2,tspc,td,alpht
      equ,3,alphr,equ,6,ctrp,const,0.0,end
difbcl=sconc,dspc,td,conc,const,3.1622e18,end
difbcr=sconc,dspc,td,conc,const,0.0,end
surfa=1.0,end
end of diffusion input
$
```

```

$ -----
equation input
$ -----
$ (1)
y=1.0,end
$ (2)
y=0.0,end
$ (3)
y=1.0e15,end
$ (4)
y=1.0e13*exp(-100./temp),end
$ (5)
y=1.0e13*exp(-500./temp),end
$ (6)
y=1.0e13*exp(-800./temp),end
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
time=0.0,end
tstep=0.01,end
timend=50.,end
nprint=500,end          $ print every 5 seconds
itermx=2000,end
delcmx=1.0e-4,end
bump=1.e-2,end
bound=4.0,end
omega=0.3,end
end of control input
$
$ -----
plot input
$ -----
nplot=100,end           $ makes plotfile entry every 1 sec
plotseg=1,end           $ segments for which plot info is needed
plotencl=end            $ no enclosure plot info is needed
dname=td,end            $ diffusing species for which plot info is needed
ename=end               $ no enclosure species plot info needed
dplot=sflux,end
eplot=end               $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1e: Diffusion with Composite Material Layers (Val-1e, see p. 15)

```
title input
  Validation Problem #5 - Tritium diffusion through PyC/SiC layer in NPR
  fuel particles at 2100 C with constant source and no trapping.
end of title input
$
$ -----
main input
$ -----
dspcnme=td,end
espcnme=t,end
segnds=13,24,end
nbrencl=2,end
linksegs=1,2,end
end of main input
$
$ -----
enclosure input
$ -----
start bdry,1,end
etemp=2373.0,end
esppres=t,1.e6,end
$
$ -----
start bdry,2,end
$ -----
etemp=2373.0,end
esppres,t,0.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
start thermseg,end
delx=0.0,11*3.0e-6,0.0,end
tempd=13*2373.0,end
$
start thermseg,end
delx=0.0,22*3.0e-6,0.0,end
tempd=24*2373.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
start diffseg,end
$ [DIFFSEG 1] PyC
nbrden=4.8319e28,end
concd=td,13*0.0,end
qstrdr=td,const,0.0,end          $ Q*/R for Soret effect unknown
dcoef=td,equ,1,end              $ Diffusion coeff (m2/s)
srcsd=td,const,0.0,srcpf,13*0.0,end
difbcl=sconc,dspc,td,conc,const,3.0537e25,end
difbcr=link,td,solcon,equ,3,end
surfa=2.16e-6,end
```

```

$
start diffseg,end
$ [DIFFSEG 2] SiC
concd=td,24*0.0,end
dcoef=td,equ,2,end
qstrdr=td,const,0.0,end
srcsd=td,const,0.0,srcpf,24*0.0,end
difbcr=sconc,dspc,td,conc,const,0.0,end
difbcl=link,td,solcon,equ,3,end
surfa=2.16e-6,end
end of diffusion input
$
$ -----
equation input
$ -----
$ (1) Diffusion coefficient PyC
y=1.274e-7,end
$ (2) Diffusion coefficient SiC
y=2.622e-11,end
$ (3) Solubility
y=1.0,end
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
time=0.0,end
tstep=0.001,end
timend=50.0,end
nprint=1000,end          $ print every second
itermx=2000,end
delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
nplot=1000,end           $ makes plotfile entry every 1 sec
plotseg=1,end            $ segments for which plot info is needed
plotencl=1,2,end         $ enclosures for which plot info is needed
dname=td,end             $ diffusing species for which plot info is needed
ename=t,end              $ enclosure species for which plot info is needed
dplot=moblinv,end
eplot=diff,end           $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Problem 1f: Heat Sink/Source Problem (Val-1fa, see p. 18)

```
title input
  Validation Problem #6a - Model Utilizes TMAP7 Thermal Capabilities
  Head Conduction in Slab with Internal Heat Generation
end of title input
$
$ -----
main input
$ -----
  dspcnme=qd,end
  espcnme=q,end
  segnds=18,end
  nbrencl=1,end
end of main input
$
$ -----
enclosure input
$ -----
  start bdry,1
  etemp=300.0,end
  espPRES=q,0.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,16*0.10,0.0,end
  tempd=18*1000.0,end
  tcon=const,10.0,end
  rhocp=const,1.0,end
  hsrc=const,1.0e4,srcpf,0.0,16*1.0,0.0,end
  htrbcl=adiab,end
  htrbcr=stemp,const,300.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0,end
  concd=qd,18*0.0,end
  dcoef=qd,const,0.1,end
  qstrdr=qd,const,0.0,end
  srcsd=qd,const,0.0,srcpf,18*0.0,end
  difbcl=nonflow,end
  difbcr=sconc,dspc,qd,conc,const,0.0,end
  surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
end of equation input
$
```

```

$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
    time=0.0,end
    tstep=0.001,end
    timend=50.1,end
    nprint=10000,end
    itermx=200,end
    delcmx=1.0e-6,end
    bump=1.e-2,end
    bound=2.0,end
    omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
    nplot=1000,end
    plotseg=end
    plotencl=1,end
    dname=end
    ename=q,end
    dplot=end
    eplot=etemp,end
end of plot input
$
end of data

```

Problem 1fb. Thermal Diffusion Transient (Val-1fb, see p. 19)

```
title input
  Validation Problem #6b - Model Utilizes TMAP7 Thermal Capabilities
  Prediction of slab Temperature as a function of time
end of title input
$
$ -----
main input
$ -----
  dspcnme=td,end
  espcnme=t,end
  segnds=18,end
  nbrencl=1,end
end of main input
$
$ -----
enclosure input
$ -----
  start bdry,1
  etemp=373.0,end
  espres=t,0.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,16*2.5e-1,0.0,end
  tempd=18*300.0,end
  tcon=const,100.0,end
  rhocp=const,100.0,end
  hsrc=const,0.0,srcpf,18*0.0,end
  htrbcl=stemp,const,400.0,end
  htrbcr=stemp,const,300.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0,end
  concd=td,18*0.0,end
  dcoef=td,const,1.0,end
  qstrdr=td,const,0.0,end
  srcsd=td,const,0.0,srcpf,18*0.0,end
  difbcl=nonflow,end
  difbcr=nonflow,end
  surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
end of equation input
$
```



```

$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
    time=0.0,end
    tstep=0.01,end
    timend=5.0,end
    nprint=10,end
    itermx=2000,end
    delcmx=1.0e-6,end
    bump=1.e-2,end
    bound=2.0,end
    omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
    nplot=10,end
    plotseg=1,end
    plotencl=1,end
    dname=td,end
    ename=t,end
    dplot=sconc,end
    eplot=end
end of plot input
$
end of data

```

Conduction in Composite Structure with Constant Surface Temperatures (Val-1fc, see p. 20)

```
title input
  Validation Problem #6c - Model Utilizes TMAP7 Thermal Capabilities
  Prediction of Composite Slab Temperature as a Function of Time
end of title input
$
$
main input
  dspcnme=td,end
  espcnme=t,end
  segnds=22,22,end
  nbrencl=2,end
  linksegs=1,2,end
end of main input
$
$
enclosure input
  start bdry,1
    etemp=600.0,end
    esppres=t,0.0,end
  start bdry,2
    etemp=600.0,end
    esppres=t,0.0,end
end of enclosure input
$
$
thermal input
  start thermseg,end
    delx=0.0,20*2.0e-2,0.0,end
    tempd=22*0.0,end
    tcon=const,401.0,end
    rhocp=const,3.4392e6,end
    hsrc=const,0.0,srcpf,22*0.0,end
    htrbcl=stemp,const,600.0,end
    htrbcr=link,end
    hgap=const,1.0e8,end
  start thermseg,end
    delx=0.0,20*2.0e-2,0.0,end
    tempd=22*0.0,end
    tcon=const,80.2,end
    rhocp=const,3.5179e6,end
    hsrc=const,0.0,srcpf,22*0.0,end
    htrbcl=link,end
    htrbcr=stemp,const,0.0,end
end of thermal input
$
$
diffusion input
  start diffseg,end
    nbrden=1.0,end
    concd=td,22*0.0,end
    dcoef=td,const,117.0e-6,end
    qstrdr=td,const,0.0,end
    srcsd=td,const,0.0,srcpf,22*0.0,end
    difbcl=sconcd,dspc,td,concd,const,600.0,end
```

```

        difbcr=link,td,solcon,const,1.0,end
        surfa=1.0,end
    start diffseg,end
        nbrden=1.0,end
        concd=td,22*0.0,end
        dcoef=td,const,23.1e-6,end
        qstrdr=td,const,0.0,end
        srcsd=td,const,0.0,srcpf,22*0.0,end
        difbcr=sconc,dspc,td,conc,const,0.0,end
        difbcl=link,td,solcon,const,1.0,end
        surfa=1.0,end
end of diffusion input
$
$
equation input
end of equation input
$
$
table input
end of table input
$
$
control input
    time=0.0,end
    tstep=0.005,end
    timend=150.005,end
    nprint=1000,end
    itermx=2000,end
    delcmx=1.0e-6,end
    bump=1.e-2,end
    bound=2.0,end
    omega=1.3,end
end of control input
$
$
plot input
    nplot=1,end
    plotseg=1,end
    plotencl=1,end
    dname=td,end
    ename=t,end
    dplot=sconc,end
    eplot=end
end of plot input
$
end of data

```

Heat Conduction in Semi-Infinite Copper Slab with Convection (Val-1fd, see p. 24)

```
title input
  Validation Problem #6d - Model Utilizes TMAP7 Thermal Capabilities
  Heat Conduction in Semi-Infinite Copper Slab with Convection
end of title input
$
$
main input
  dspcnme=qd,end
  espcnme=q,end
  segnds=90,end
  nbrencl=1,end
end of main input
$
$
enclosure input
  start bdry,1
  etemp=500.0,end
  esppres=q,0.0,end
end of enclosure input
$
$
thermal input
  start thermseg,end
  delx=0.0,16*0.1,0.5,1.0,70*5.0,0.0,end
  tempd=90*100.0,end
  tcon=const,401.0,end
  rhocp=const,3.439e6,end
  hsrc=const,0.0,srcpf,0.0,88*0.0,0.0,end
  htrbcl=convec,const,200.0,encl,1,end
  htrbcr=stemp,const,0.0,end
end of thermal input
$
$
diffusion input
  start diffseg,end
  nbrden=1.0,end
  concd=qd,90*0.0,end
  dcoef=qd,const,0.1,end
  qstrdr=qd,const,0.0,end
  srcsd=qd,const,0.0,srcpf,90*0.0,end
  difbcl=nonflow,end
  difbcr=sconc,dspc,qd,conc,const,0.0,end
  surfa=1.0,end
end of diffusion input
$
$
equation input
end of equation input
$
$
table input
end of table input
$
$
```

```

control input
  time=0.0,end
  timestep=0.01,end
  timend=1290.01,end
  nprint=1000,end
  itermx=200,end
  delcmx=1.0e-6,end
  bump=1.e-2,end
  bound=2.0,end
  omega=1.3,end
end of control input
$
$
plot input
  nplot=1000,end
  plotseg=end
  plotencl=1,end
  dname=qd,end
  ename=q,end
  dplot=end
  eplot=etemp,end
end of plot input
$
end of data

```

Problem 1ga. Simple Forward Reactions (Val-1ga, see p.26)

```
title input
  Validation Problem #7a - Simple Chemical Reaction Problem
  Equal Starting Concentrations
end of title input
$
$ -----
main input
$ -----
  dspcnme=q,end
  espcnme=a,b,ab,end
  segnds=3,end
  nbrencl=1,end
end of main input
$
$ -----
enclosure input
$ -----
  start func,1
  etemp=300.0,end
  espres=a,1.0e-6,b,1.0e-6,ab,0.0,end
  reaction=nequ,1
    ratequ,1
    nreact,2,a,1.0,b,1.0
    nprod,1,ab,1.0,end
  evol=10.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,1.0,0.0,end
  tempd=3*300.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0,end
  concd=q,3*0.0,end
  dcoef=q,const,1.0,end
  qstrdr=q,const,0.0,end
  srcsd=q,const,0.0,srcpf,3*0.0,end
  difbcl=nonflow,end
  difbcr=nonflow,end
  surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
  y=4.14e-15*conce(1)*conce(2),end
end of equation input
```

```

$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
  time=0.0,end
  tstep=0.01,end
  timend=50.0,end
  nprint=1000,end
  itermx=200,end
  delcmx=1.0e-6,end
  bump=1.e-2,end
  bound=2.0,end
  omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
  nplot=100,end
  plotseg=end
  plotencl=1,end
  dname=end
  ename=a,b,ab,end
  dplot=end
  eplot=press,end
end of plot input
$
end of data

```

Problem 1gb. Simple Forward Reactions (Val-1gb, see p. 26)

```
title input
  Validation Problem #7b - Simple Chemical Reaction Problem
  Unequal Starting Concentrations
end of title input
$
$ -----
main input
$ -----
  dspcnme=q,end
  espcnme=a,b,ab,end
  segnds=3,end
  nbrencl=1,end
end of main input
$
$ -----
enclosure input
$ -----
  start func,1
  etemp=300.0,end
  espres=a,1.0e-6,b,5.0e-7,ab,0.0,end
  reaction=nequ,1
    ratequ,1
    nreact,2,a,1.0,b,1.0
    nprod,1,ab,1.0,end
  evol=10.0,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,1.0,0.0,end
  tempd=3*300.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0,end
  concd=q,3*0.0,end
  dcoef=q,const,1.0,end
  qstrdr=q,const,0.0,end
  srcsd=q,const,0.0,srcpf,3*0.0,end
  difbcl=nonflow,end
  difbcr=nonflow,end
  surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
  y=4.14e-15*conce(1)*conce(2),end
end of equation input
```



```

$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
  time=0.0,end
  tstep=0.001,end
  timend=50.0,end
  nprint=10000,end
  itermx=200,end
  delcmx=1.0e-6,end
  bump=1.e-2,end
  bound=2.0,end
  omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
  nplot=1000,end
  plotseg=end
  plotencl=1,end
  dname=end
  ename=a,b,ab,end
  dplot=end
  eplot=press,end
end of plot input
$
end of data

```

Problem 1gc. Series Reactions (Val-gc, see p. 29)

```
title input
  Validation Problem #7c - Chemical Reaction in Series Problem
  a -> b -> c
end of title input
$
$ -----
main input
$ -----
  dspcnme=q,end
  espcnme=a,b,c,end
  segnds=3,end
  nbrencl=1,end
end of main input
$
$ -----
enclosure input
$ -----
  start func,1
  etemp=300.0,end
  espres=a,1.0e-6,b,0.0,c,0.0,end
  reaction=nequ,2
    ratequ,1
    nreact,1,a,1.0,nprod,1,b,1.0
    ratequ,2
    nreact,1,b,1.0,nprod,1,c,1.0,end
  evol=1.5e-1,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,1.0,0.0,end
  tempd=3*300.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0,end
  concd=q,3*0.0,end
  dcoef=q,const,1.0,end
  qstrdr=q,const,0.0,end
  srcsd=q,const,0.0,srcpf,3*0.0,end
  difbcl=nonflow,end
  difbcr=nonflow,end
  surfa=1.0,end
end of diffusion input
$
$ -----
equation input
$ -----
$ (1)
```

```

y=1.25e-2*conce(1),end
$ (2)
y=2.5e-3*conce(2),end
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
    time=0.0,end
    tstep=0.01,end
    timend=901.0,end
    nprint=20,end
    itermx=200,end
    delcmx=1.0e-6,end
    bump=1.e-2,end
    bound=2.0,end
    omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
    nplot=100,end
    plotseg=end
    plotencl=1,end
    dname=end
    ename=a,b,c,end
    dplot=end
    eplot=press,end
end of plot input
$
end of data

```

Problem 1ha. Three Enclosure Problem (Val-1ha, See p. 32)

```
title input
  Validation Problem #8a - System (Multiple Enclosure Volumes) Problem
end of title input
$
$ -----
main input
$ -----
  dspcnme=t,end
  espcnme=t2,end
  segnds=3,end
  nbrencl=3,end
end of main input
$
$ -----
enclosure input
$ -----
  start bdry,1
    etemp=303.,end
    espres=t2,const,1.0,end
    outflow = nbrflwp,1,qflow,const,0.1,rencl,2,end
  start func,2
    etemp=303.0,end
    espres=t2,0.0,end
    evol=1.0,end
    outflow=nbrflwp,1,qflow,const,0.1,rencl,3,end
  start func,3
    etemp=303.0,end
    espres=t2,0.0,end
    evol = 1.0,end
    outflow=nbrflwp,1,qflow,const,0.1,rencl,1,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,1.0,0.0,end
  tempd=3*303.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0e21,end
  concd=t,3*0.0,end
  dcoef=t,const,1.0,end
  qstrdr=t,const,0.0,end
  srcsd=t,const,0.0,srcpf,3*0.0,end
  difbcl=nonflow,end
  difbcr=nonflow,end
  surfa=1.0,end
end of diffusion input
$
```

```

$ -----
equation input
$ -----
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
  time=0.0,end
  tstep=0.001,end
  timend=40.001,end
  nprint=10000,end
  itermx=200,end
  delcmx=1.0e-6,end
  bump=1.e-2,end
  bound=2.0,end
  omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
  nplot=1000,end
  plotseg=end
  plotencl=2,3,end
  dname=end
  ename=t2,end
  dplot=end
  eplot=conv,press,end
end of plot input
$
end of data

```

Problem 1hb. Equilibrating Enclosures (Val-1hb, see p. 33)

```

title input
  Validation Problem #8b - System Problem with Different Starting
  pressures
end of title input
$
$ -----
main input
$ -----
  dspcnme=t,end
  espcnme=t2,d2,end
  segnds=3,end
  nbrencl=2,end
end of main input
$
$ -----
enclosure input
$ -----
  start func,1
    etemp=303.0,end
    espmpres=t2,1.0,d2,0.0,end
    reaction=nequ,0,end
    evol=1.0,end
    outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
  start func,2
    etemp=303.0,end
    reaction = nequ,0,end
    espmpres=t2,0.0,d2,1.0,end
    evol=1.0,end
    outflow=nbrflwp,1,qflow,const,0.1,rencl,1,end
end of enclosure input
$
$ -----
thermal input
$ -----
  start thermseg,end
  delx=0.0,1.0,0.0,end
  tempd=3*303.0,end
end of thermal input
$
$ -----
diffusion input
$ -----
  start diffseg,end
  nbrden=1.0e21,end
  concd=t,3*0.0,end
  dcoef=t,const,1.0,end
  qstrdr=t,const,0.0,end
  srcsd=t,const,0.0,srcpf,3*0.0,end
  difbcl=nonflow,end
  difbcr=nonflow,end
  surfa=1.0,end
end of diffusion input
$
$ -----

```

```

equation input
$ -----
end of equation input
$
$ -----
table input
$ -----
end of table input
$
$ -----
control input
$ -----
time=0.0,end
tstep=0.001,end
timend=40.001,end
nprint=10000,end
itermx=200,end
delcmx=1.0e-6,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
end of control input
$
$ -----
plot input
$ -----
nplot=1000,end
plotseg=end
plotencl=1,2,end
dname=end
ename=t2,d2,end
dplot=end
eplot=conv,press,end
end of plot input
$
end of data

```

Species Equilibration on a Reactive Surface (Val-1ia, see p. 37)

```

title input
Problem #9a. Chemical equilibration on polycrystalline tungsten surface
using conventional dissociation-recombination boundary condition.
end of title input
$ -----
main input
$ -----
dspcnme=h,d,end
espcnme=h2,d2,hd,end
segnds=12,end
nbrencl=1,end          $ test chamber
end of main input
$
$ -----
enclosure input
$ -----
start func,1,end          $ Test chamber where sample is
$ Enclosure 1 is the test chamber with equal starting pressures
etemp=const,1000.0,end
esppres=h2,1.0e4,d2,1.0e4,hd,1.0e-10,end
evol=1.0,end          $ Assumed value of 1.0 m3
end of enclosure input
$
thermal input
start thermseg,end
$ 1-mm foil [THERMSEG 1]
delx=0.0,10*1.0e-4,0.0,end
tempd=12*1000.,end          $ Constant temperature (K)
end of thermal input
$ =====
diffusion input
$ =====
start diffseg,end
$ 1-mm foil [DIFFSEG 1]
nbrden=6.25e28,end
concd=h,const,1.0,d,const,1.0,end  $ Starting mobile concentrations
qstrdr=h,const,0.0,d,const,0.0,end  $ Q*/R for Soret effect unknown
dcoef=h,eq,1,d,eq,1,end
srcsd=h,const,0.0,srcpf,const,1.0,d,const,0.0,srcpf,const,1.0,end
difbcl=ratedep,encl,1,
      spc,h,exch,h2,ksubd,eq,2,h,ksubr,1.29e-16
      exch,hd,ksubd,eq,2,d,ksubr,2.58e-16
      spc,d,exch,hd,ksubd,eq,2,h,ksubr,2.58e-16
      exch,d2,ksubd,eq,2,d,ksubr,1.29e-16,end
difbcr=nonflow,end
surfa=0.0025,end          $ 50 x 50 mm square
end of diffusion input
$
equation input
$ (1) Diffusivity for h in tungsten (m2/s)
y=4.1e-7*exp(-3.39/8.625e-5/temp),end  $modified from 0.39 eV
$ (2) Dissociation coefficient at full efficiency
y=1.85802e24/sqrt(temp),end
end of equation input

```



```

$
table input
end of table input
$
control input
time=0.,end
tstep=0.01,end
timend=6.1,end
nprint=100,end
itermx=15000,end
delcmx=1.e-6,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
damp=0.7,end
end of control input
$
plot input
nplot=20,end          $ makes plotfile entry every 0.2 sec
plotseg=1,end         $ segments for which plot info is needed
plotenc1=1,end        $ enclosures for which plot info is needed
dname=h,d,end         $ diffusing species for which plot info is needed
ename=h2,d2,hd,end    $ enclosure species for which plot info is needed
dplot=moblinv,end
eplot=press,diff,end
end of plot input
$
end of data

```

Species Equilibration on a Reactive Surface (Val-1ib, see p. 39)

```

title input
Problem #9b. Chemical equilibration on polycrystalline tungsten surface
using conventional dissociation-recombination boundary condition.
end of title input
$ -----
main input
$ -----
dspcnme=h,d,end
espcnme=h2,d2,hd,end
segnds=12,end
nbrencl=1,end          $ test chamber
end of main input
$
$ -----
enclosure input
$ -----
start func,1,end          $ Test chamber where sample is
$ Enclosure 1 is the test chamber with equal starting pressures
etemp=const,1000.0,end
esppres=h2,1.0e4,d2,1.0e5,hd,1.0e-10,end
evol=1.0,end          $ Assumed value of 1.0 m3
end of enclosure input
$
thermal input
start thermseg,end
$ 1-mm foil [THERMSEG 1]
delx=0.0,10*1.0e-4,0.0,end
tempd=12*1000.,end          $ Constant temperature (K)
end of thermal input
$ =====
diffusion input
$ =====
start diffseg,end
$ 1-mm foil [DIFFSEG 1]
nbrden=6.25e28,end
concd=h,const,1.0,d,const,1.0,end  $ Starting mobile concentrations
qstrdr=h,const,0.0,d,const,0.0,end  $ Q*/R for Soret effect unknown
dcoef=h,eq,1,d,eq,1,end
srcsd=h,const,0.0,srcpf,const,1.0,d,const,0.0,srcpf,const,1.0,end
difbcr=ratedep,encl,1,
      spc,h,exch,h2,ksubd,eq,2,h,ksubr,1.29e-16
      exch,hd,ksubd,eq,2,d,ksubr,2.58e-16
      spc,d,exch,hd,ksubd,eq,2,h,ksubr,2.58e-16
      exch,d2,ksubd,eq,2,d,ksubr,1.29e-16,end
difbcl=nonflow,end
surfa=0.0025,end          $ 50 x 50 mm square
end of diffusion input
$
equation input
$ (1) Diffusivity for h in tungsten (m2/s)
y=4.1e-7*exp(-3.39/8.625e-5/temp),end  $modified from 0.39 eV
$ (2) Dissociation coefficient at full efficiency
y=1.85802e24/sqrt(temp),end
end of equation input

```

```

$
table input
end of table input
$
control input
time=0.,end
tstep=0.01,end
timend=6.1,end
nprint=100,end
itermx=1500,end
delcmx=1.e-6,end
bump=1.e-2,end
bound=2.0,end
omega=1.3,end
damp=0.7,end
end of control input
$
plot input
nplot=20,end          $ makes plotfile entry every 0.2 sec
plotseg=1,end         $ segments for which plot info is needed
plotenc1=1,end        $ enclosures for which plot info is needed
dname=h,d,end         $ diffusing species for which plot info is needed
ename=h2,d2,hd,end    $ enclosure species for which plot info is needed
dplot=moblinv,end
eplot=press,diff,end
end of plot input
$
end of data

```

Species Equilibration on a Reactive Surface (Val-1ic, see p. 39)

```

title input
Problem #9. Chemical equilibration on polycrystalline tungsten surface.
end of title input
$ -----
main input
$ -----
dspcnme=h,d,end
espcnme=h2g,d2g,hdg,end
sspcnme=h2,d2,hd,end
segnds=7,end
nbrencl=1,end          $ test chamber
end of main input
$
enclosure input
start func,1,end          $ Test chamber where sample is
$ Enclosure 1 is the test chamber with equal starting pressures
etemp=const,1000.0,end
esppres=h2g,1.0e4,d2g,1.0e4,hdg,1.e-10,end
evol=1.0,end          $ Assumed value of 1.0 m3
end of enclosure input
$
thermal input
start thermseg,end
$ 1-mm foil [THERMSEG 1]
delx=0.0,5*2.0e-4,0.0,end
tempd=7*1000.,end          $ Constant temperature (K)
end of thermal input
$ =====
diffusion input
$ =====
start diffseg,end
$ 1-mm foil [DIFFSEG 1]
nbrden=6.25e28,end
concd=h,const,1.0e0,d,const,1.0e0,end  $ Starting mobile concentrations
ssconc=h2,1.0,1.0,d2,1.0,1.0,hd,1.0,1.0,end
qstrdr=h,const,0.0,d,const,0.0,end  $ Q*/R for Soret effect unknown
dcoef=h,eq,1,d,eq,1,h2,eq,1,d2,eq,1,hd,eq,1,end
srcsd=h,const,0.0,srcpf,const,1.0,d,const,0.0,srcpf,const,1.0,end
difbcl=surfdep,encl,1
    spc,h,nu,8.4e12,ec,-0.05,es,1.04
        comb,h,prob,1.0
        comb,d,prob,1.0
    spc,d,nu,8.4e12,ec,-0.05,es,1.04
        comb,h,prob,1.0
        comb,d,prob,1.0
    spc,h2,nu,8.4e12,ec,-0.05
        exch,h2g,amu,2.0,ex,0.05
        diss,h,h,eb,0.0
        form,h,h,prob,1.0
    spc,d2,nu,8.4e12,ec,-0.05
        exch,d2g,amu,2.0,ex,0.05
        diss,d,d,eb,0.0
        form,d,d,prob,1.0
    spc,hd,nu,8.4e12,ec,-0.05

```

```

        exch,hdg,amu,2.0,ex,0.05
        diss,h,d,eb,0.0
        form,h,d,prob,1.0,end
difbcr=nonflow,end
surfa=0.0025,end          $ 50 x 50 mm square
end of diffusion input
$
equation input
$ (1) Diffusivity for h,d in tungsten (m2/s)
y=5.33e-7*exp(-0.39/8.625e-5/temp),end
end of equation input
$
table input
end of table input
$
control input
time=0.,end
tstep=0.01,end
timend=10.,end
nprint=1,end
itermx=1000,end
delcmx=1.e-7,end
bump=1.e-4,end
bound=10.,end
omega=1.0,end
damp=0.7
end of control input
$
plot input
nplot=50,end              $ makes plotfile entry every 0.2 sec
plotseg=1,end             $ segments for which plot info is needed
plotencl=1,end            $ enclosures for which plot info is needed
dname=h,d,end             $ diffusing species for which plot info is needed
sname=h2,d2,hd,end        $ surface species for which plot info is needed
ename=h2g,d2g,hdg,end     $ enclosure species for which plot info is needed
dplot=moblinv,end
eplot=press,diff,end
end of plot input
$
end of data

```

Species Equilibration on a Reactive Surface (Val-1id, see p. 39)

```

title input
Problem #9d. Chemical equilibration on polycrystalline tungsten surface.
Surfdep conditions with unequal starting pressures.
end of title input
$ -----
main input
$ -----
dspcnme=h,d,end
espcnme=h2g,d2g,hdg,end
sspcnme=h2,d2,hd,end
segnds=7,end
nbrencl=1,end          $ test chamber
end of main input
$
enclosure input
start func,1,end          $ Test chamber where sample is
$ Enclosure 1 is the test chamber with equal starting pressures
etemp=const,1000.0,end
esppres=h2g,1.0e4,d2g,1.0e5,hdg,1.e-10,end
evol=1.0,end          $ Assumed value of 1.0 m3
end of enclosure input
$
thermal input
start thermseg,end
$ 1-mm foil [THERMSEG 1]
delx=0.0,5*2.0e-4,0.0,end
tempd=7*1000.,end          $ Constant temperature (K)
end of thermal input
$ =====
diffusion input
$ =====
start diffseg,end
$ 1-mm foil [DIFFSEG 1]
nbrden=6.25e28,end
concd=h,const,0.0e00,d,const,0.0e0,end  $ Starting mobile concentrations
ssconc=h2,1.0,1.0,d2,1.0,1.0,hd,1.0,1.0,end
qstrdr=h,const,0.0,d,const,0.0,end  $ Q*/R for Soret effect unknown
dcoef=h,eq,1,d,eq,1,h2,eq,1,d2,eq,1,hd,eq,1,end
srcsd=h,const,0.0,srcpf,const,1.0,d,const,0.0,srcpf,const,1.0,end
difbcl=surfdep,encl,1
    spc,h,nu,8.4e12,ec,-0.01,es,1.04
        comb,h,prob,1.0
        comb,d,prob,1.0
    spc,d,nu,8.4e12,ec,-0.01,es,1.04
        comb,h,prob,1.0
        comb,d,prob,1.0
    spc,h2,nu,8.4e12,ec,-0.01
        exch,h2g,amu,2.0,ex,0.05
        diss,h,h,eb,0.0
        form,h,h,prob,1.0
    spc,d2,nu,8.4e12,ec,-0.01
        exch,d2g,amu,2.0,ex,0.05
        diss,d,d,eb,0.0
        form,d,d,prob,1.0

```

```

    spc,hd,nu,8.4e12,ec,-0.01
    exch,hdg,amu,2.0,ex,0.05
    diss,h,d,eb,0.0
    form,h,d,prob,1.0,end
difbcr=nonflow,end
surfa=0.0025,end          $ 50 x 50 mm square
end of diffusion input
$
equation input
$ (1) Diffusivity for h,d in tungsten (m2/s)
y=5.33e-7*exp(-0.39/8.625e-5/temp),end
end of equation input
$
table input
end of table input
$
control input
time=0.,end
tstep=0.01,end
timend=10.,end
nprint=100,end
itermx=19000,end
delcmx=1.e-6,end
bump=1.e-4,end
bound=1.1,end
omega=1.3,end
damp=0.7
end of control input
$
plot input
nplot=50,end              $ makes plotfile entry every 0.2 sec
plotseg=1,end             $ segments for which plot info is needed
plotencl=1,end            $ enclosures for which plot info is needed
dname=h,d,end             $ diffusing species for which plot info is needed
sname=h2,d2,hd,end        $ surface species for which plot info is needed
ename=h2g,d2g,hdg,end    $ enclosure species for which plot info is needed
dplot=moblinv,end
eplot=press,diff,end
end of plot input
$
end of data

```

Problem 1ja: Radioactive Decay of Mobile Tritium in a Slab (Val-1ja, see p. 41)

```
title input
  Validation Problem #10a -    1st order decay in lattice
    T -- >    He-3
end of title input
$
main input
dspcnme=t,he,end
dkrate=t,1.782411e-9,he,end
espcnme=ts,end
segnds=17,end
nbrencl=1,end
end of main input
$
enclosure input
start bdry,1,end
etemp=273.0,end
esppres=ts,1.0e5,end
end of enclosure input
$
thermal input
start thermseg,end
delx=0.0,15*0.1,0.0,end
tempd=17*273.0,end
end of thermal input
$
diffusion input
start diffseg,end
nbrden=4.832e28,end
concd=t,17*1.0,he,17*0.0,end
qstrdr=t,const,0.0,he,const,0.0,end
dcoef=t,eq,1,he,const,0.0,end
srcsd=t,const,0.0,srcpf,17*0.0,he,const,0.0,srcpf,17*0.0,end
difbcl=nonflow,end
difbcr=nonflow,end
surfa=1.0,end
end of diffusion input
$
equation input
$ (1) Diffusivity for t
y=1.58e-4*exp(-308000.0/(8.314*temp)),end
$ (2) Trap release frequency
y=1.0e13*exp(-4.2/8.124e-5/temp),end
$ (3) Trapping frequency for t
y=2.096e15*exp(-308000.0/(8.314*temp)),end
end of equation input
$
table input
end of table input
$
control input
time=0.0,end
tstep=1.15e5,end           $ time step = .01 year
timend=1.4197e9,end        $ 45 years
nprint=100,end             $ print every year
```



```

itermx=20000,end
delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
damp=0.7,end
omega=1.3,end
end of control input
$
plot input
nplot=100,end          $ makes plotfile entry every 1/10 year
plotseg=1,end          $ segments for which plot info is needed
plotencl=end           $ enclosure info is not needed
dname=t,he,end         $ diffusing species for which plot info is needed
ename=end              $ enclosure species for which plot info is needed
dplot=moblinv,trapinv,end
eplot=end
end of plot input
$
end of data

```

Problem 1jb: Decay of Tritium in a Distributed Trap (Val-1jb, see p. 43)

```
title input
  Validation Problem #10b -    1st order decay in traps
    T -- >    He-3
end of title input
$
main input
dspcnme=t,he,end
dkrate=t,1.782411e-9,he,end
espcnme=ts,end
segnds=17,end
nbrencl=1,end
end of main input
$
enclosure input
start bdry,1,end
etemp=273.0,end
esppres=ts,1.0e5,end
end of enclosure input
$
thermal input
start thermseg,end
delx=0.0,15*0.1,0.0,end
tempd=17*273.0,end
end of thermal input
$
diffusion input
start diffseg,end
nbrden=4.832e28,end
concd=t,17*1.0,he,17*0.0,end
qstrdr=t,const,0.0,he,const,0.0,end
dcoef=t,eq,1,he,const,0.0,end
srcsd=t,const,0.0,srcpf,17*0.0,he,const,0.0,srcpf,17*0.0,end
difbcl=nonflow,end
difbcr=nonflow,end
trapping=ttyp,1,tconc,norm,0.001,0.75,0.5,0.0,tspc,t,alphr,eq,2
      alphr,eq,3,ctrap,const,0.5,end
surfa=1.0,end
end of diffusion input
$
equation input
$ (1) Diffusivity for t
y=1.58e-4*exp(-308000.0/(8.314*temp)),end
$ (2) Trap release frequency
y=1.0e13*exp(-4.2/8.124e-5/temp),end
$ (3) Trapping frequency for t
y=2.096e15*exp(-308000.0/(8.314*temp)),end
end of equation input
$
table input
end of table input
$
control input
time=0.0,end
tstep=1.15e5,end                                $  time step = .01 year
```

```

timend=1.4197e9,end          $ 45 years
nprint=100,end              $ print every year
itermx=20000,end
delcmx=1.0e-7,end
bump=1.e-2,end
bound=2.0,end
damp=0.7,end
omega=1.3,end
end of control input
$
plot input
nplot=100,end               $ makes plotfile entry every 1/10 year
plotseg=1,end              $ segments for which plot info is needed
plotencl=end               $ enclosure info is not needed
dname=t,he,end             $ diffusing species for which plot info is needed
ename=end                  $ enclosure species for which plot info is needed
dplot=moblinv,trapinv,end
eplot=end
end of plot input
$
end of data

```

Ion Implantation Experiment (Val-2a, see p. 45)

```
title input
  Sample Problem #1 - Plasma driven premeation of PCA
end of title input
$
$
main input
  dspcnme=d,end
  espcnme=d2,end
  segnds=21,end
  nbrencl=2,end
end of main input
$
$
enclosure input
  start bdry,1,end
  etemp=703.,end
  espPRES=d2,tabl,1,end
$
  start bdry,2,end
  etemp=703.0,end
  espPRES=d2,const,2.e-6,end
end of enclosure input
$
$
thermal input
  start thermseg,end
  delx=0.0,5*4.0e-9,1.0e-8,1.0e-7,1.0e-6
  1.0e-5,10*4.88e-5,0.0,end
  tempd=21*703.0,end
end of thermal input
$
$
diffusion input
  start diffseg,end
  nbrden=6.45e28,end
  concd=d,21*0.0,end
  dcoef=d,const,3.0e-10,end
  qstrdr=d,const,0.0,end
  srcsd=d,tabl,2,srcpf,3*0.0,0.25,1.0,0.25,15*0.0,end
  difbcl=ratedep,encl,1,spc,d
  exch,d2,ksubd,equ,1,d,ksubr,equ,2,end
  difbcr=ratedep,encl,2,spc,d
  exch,d2,ksubd,const,1.7918e15,d,ksubr
  const,2.0e-31,end
  surfa=1.0,end
end of diffusion input
$
$
equation input
$ (1) Dissociation constant ( $d_2/M^2.s.Pa^{1/2}$ )
y= 8.959e18*(1.0-0.9999*exp(-6.0e-5*time)),end
$ (2) Recombination constant ( $m^4/d_2.s$ )
y= 1.0e-27*(1.0-0.9999*exp(-6.0e-5*time)),end
end of equation input
```

```

$
$
table input
$ (1) Upstream enclosure pressure history
0.0,4.0e-5,6420.0,4.0e-5,6420.1,9.0e-6,9420.0,9.0e-6,9420.1,4.0e-5
12480.0,4.0e-5,12480.1,9.0e-6,14940.0,1.9e-6,14940.1,4.0e-5,18180.0
4.0e-5,18180.1,9.0e-6,1.0e10,9.0e-6,end
$ (2) Implantation Flux (d/m2.s)
0.0,4.9e19,6420.0,4.9e19,6420.1,0.0,9420.0,0.0,9420.1,4.9e19
12480.0,4.9e19,12480.1,0.0,14940.0,0.0,14940.1,4.9e19,18180.0
4.9e19,18180.1,0.0,1.0e10,0.0,end
end of table input
$
$
control input
    time=0.0,end
    tstep=2.0,end
    timend=19200.0,end
    nprint=600,end
    itermx=9000,end
    delcmx=1.0e-7,end
    bump=1.e-2,end
    bound=2.0,end
    omega=1.3,end
end of control input
$
$
plot input
    nplot=30,end
    plotseg=1,end
    plotencl=1,2,end
    dname=d,end
    ename=d2,end
    dplot=moblinv,sflux,end
    eplot=end
end of plot input
$
end of data

```

Diffusion Experiment in Beryllium (Val-2ba, Val-2bb, see p. 46)

Charging Segment

```
title input
  Sample Problem #2a - R. G. Macaulay-Newcombe's thermal charging problem for
  gas absorption into a wafer of polished beryllium with a thin oxide film.
end of title input
$
$
main input
  dspcnme=d,end
  espcnme=d2,end
  segnds=20,17,end
  nbrencl=1,end
  linksegs=1,2,end
end of main input
$
$
enclosure input
  start bdry,1,end
  etemp=773.,end
  espres=d2,equ,6,end
end of enclosure input
$
$
thermal input
$ Segment 1 - BeO film
  start thermseg,end
  delx=0.0,18*1.0e-9,0.0,end
  tempd=20*773.0,end
  tcon=const,159.2,end
  rhocp=const,3.0e6,end
  hsrc=const,0.0,srcpf,20*0.0,end
  htrbcl=stemp,equ,1,end
  htrbcr=link,end
  hgap=const,1.e6,end
$ Segment 2 - Be metal - half thick
  start themseg,end
  delx=0.0,1.0e-9,1.e-8,1.e-7,1.e-6,1.e-5,10*1.888e-5,0.0,end
  tempd=17*773.0,end
  tcon=const,168.0,end
  rhocp=const,3.37e6,end
  hsrc=const,0.0,srcpf,17*0.0,end
  htrbcl=link,end
  htrbcr=adiab,end
end of thermal input
$
$
diffusion input
$ Segment 1 - BeO flim
  start diffseg,end
  nbrden=1.238e29,end
  concd=d,20*0.0,end
  dcoef=d,equ,2,end
  qstrdr=d,const,0.0,end
  srcsd=d,const,0.0,srcpf,20*0.0,end
```

```

        difbcl=lawdep,encl,1,dspc,d,d2
            pexp,0.5,solcon,equ,3,end
        difbcr=link,d,solcon,equ,3,end
        surfa=1.04e-4,end
$ Segment 2 - Be foil - foil thickness
start diffseg,end
    nbrden=1.238e29,end
    concd=d,17*0.0,end
    dcoef=d,equ,4,end
    qstrdr=d,const,0.0,end
    srcsd=d,const,0.0,srcpf,17*0.0,end
    difbcl=link,d,solcon,equ,5,end
    difbcr=nonflow,end
    surfa=1.04e-4
end of diffusion input
$
$
equation input
$ (1) Temperature History Equation
y= 773.-int(time/180000.)*(1-exp(-(time-180000.)/2700.))*475.,end
$ (2) - (5) Diffusion and Solubility Equations
$ (2) D of d in BeO (Fowler 1)
    y= 1.40e-4*exp(-24408./temp),end
$ (3) S for d in BeO
    y=5.00e20*exp(9377.7/temp),end
$ (4) D of D in Be (Abramov Be-2)
    y=8.0e-9*exp(-4220./temp),end
$ (5) S for d in Be (Swansiger)
    y=7.156e27*exp(-11606./temp),end
$ (6) Pressure History
y=13300.0*(1-int(time/180015.))+1.0e-6,end
end of equation input
$
$
table input
end of table input
$
$
control input
    time=0.0,end
    tstep=60.0,end
    timend=182400.0,end
    nprint=90,end
    itermx=90,end
    delcmx=1.0e-8,end
    bump=1.e-2,end
    bound=2.0,end
    omega=1.3,end
end of control input
$
$
plot input
    nplot=1,end
    plotseg=1,2,end
    plotencl=end
    dname=d,end
    ename=end

```

```

    dplot=moblinv,sflux,stemp,end
    eplot=end
end of plot input
$
end of data

```

Desorption Segment (Restart)

```

restart
$
equation input
$ (1) Temperature History Equation
y= 300.0+0.05*time,end
$ (2) - (5) Diffusion and Solubility Equations
$   (2) D of d in BeO (Fowler 1)
      y= 7.00e-5*exp(-27000./temp),end
$   (3) S for d in BeO
      y=5.00e20*exp(9377.7/temp),end
$   (4) D of D in Be (Abramov Be-2)
      y=8.0e-9*exp(-4220./temp),end
$   (5) S for d in Be (Swansiger)
      y=7.156e27*exp(-11606./temp),end
$ (6) Pressure History
y=0.001,end
end of equation input
$
table input
end of table input
$
control input
  time=0.0,end
  tstep=60.0,end
  timend=15460.0,end
  nprint=10,end
  itermx=90,end
  delcmx=1.0e-8,end
  bump=1.e-2,end
  bound=2.0,end
  omega=1.3,end
end of control input
$
plot input
  nplot=10,end
  plotseg=1,2,end
  plotencl=end
  dname=d,end
  ename=end
  dplot=moblinv,sflux,end
  eplot=end
end of plot input
$
end of data

```


Test Cell Release Experiment (Val-2c, see p. 48)

```
title input
  Sample Problem #3 - HTO history in an exposure chamber at TSTA
end of title input
$
$
main input
  dspcnme=t2d,htd,htod,h2od,end
  espcnme=t2,ht,hto,h2o,end
  segnds=12,end
  nbrencl=4,end
end of main input
$
$
enclosure input
$ Room air source
  start bdry,1,end
  etemp=303.,end
  espPRES=t2,0.,ht,0.,hto,0.,h2o,714.,end
  outflow=nrflwp,2,qflow,const,1.33333e-4,rencl,2,
    qflow,const,1.66667e-5,rencl,4,end
$ Exposure chamber
  start func,2,end
  etemp=303.,end
  espPRES=t2,2.e-30,ht,1.0e-30,hto,1.0e-30,h2o,714.,end
  outflow=nrflwp,1,qflow,const,1.5e-4,rencl,3,end
  reaction=nequ,2,ratequ,1
    nreact,2,t2,1.,h2o,1.,nprod,2,hto,1.,ht,1.
    ratequ,2
    nreact,2,ht,1.,h2o,1.,nprod,1,hto,1.,end
  evol=0.96,end
$ Exhaust to stack
  start bdry,3,end
  etemp=303.,end
  espPRES=t2,0.0,ht,0.,hto,0.,h2o,714.,end
$ Tritium source container
  start func,4,end
  etemp=573.,end
  espPRES=t2,2737.,ht,82.,hto,1.e-4,h2o,0.0,end
  outflow=nrflwp,1,qflow,const,1.66667e-5,rencl,2,end
  evol=3.e-4,end
end of enclosure input
$
$
thermal input
  start thermseg,end
  delx=0.0,10*1.6e-5,0.0,end
  tempd=12*303.0,end
end of thermal input
$
$
diffusion input
$ Segment 1
  start diffseg,end
  nbrden=7.65e28,end
```

```

        concd=t2d,12*0.,htd,12*0.,htod,12*0.,h2od,12*0.,end
        dcoef=t2d,const,4.e-12,htd,const,4.e-12,htod,const,1.e-14,
            h2od,const,1.e-14,end
        qstrdr=t2d,const,0.,htd,const,0.,htod,const,0.,h2od,const,0.,end
        srcsd=t2d,const,0.0,srcpf,12*0.,htd,const,0.,srcpf,12*0.
            htod,const,0.,srcpf,12*0.,h2od,const,0.,srcpf,12*0.,end
        difbcl=lawdep,encl,2,dspc,t2d,t2,pexp,1.,solcon,const,4.e19
            dspc,htd,ht,pexp,1.,solcon,const,4.e19
            dspc,htod,hto,pexp,1.,solcon,const,6.e24
            dspc,h2od,h2o,pexp,1.,solcon,const,6.e24,end

        difbcr=nonflow,end
        surfa=5.6,end
end of diffusion input
$
$
equation input
$ (1) - (2) Reaction Rate Equations
$ Index forc conc array is relative enclosure specie number
$      (i.e., t2=1, ht=2, hto=3, h2o=4)
$ (1)
y= 2.0e-29*conce(1)*(2.*conce(1)+conce(2)+conce(3)),end
$ (2)
y= 1.0e-29*conce(2)*(2.*conce(1)+conce(2)+conce(3)),end
end of equation input
$
$
table input
end of table input
$
$
control input
    time=0.0,end
    tstep=6.0,end
    timend=180000.0,end
    nprint=600,end
    itermx=90,end
    delcmx=1.0e-6,end
    bump=1.e-2,end
    bound=3.0,end
    omega=1.3,end
    damp=0.7
end of control input
$
$
plot input
    nplot=50,end
    plotseg=1,end
    plotencl=2,4,end
    dname=t2d,htd,htod,htod,end
    ename=t2,ht,hto,h2o,end
    dplot=moblinv,sflux,end
    eplot=press,conv,end
end of plot input
$
end of data

```

Problem 2d. Thermal Desorption Spectroscopy on Tungsten (Val-2d, see p. 49)

```

title input
Simulation of polycrystalline tungsten experiment irradiated at RT with
H at 5 keV, 1E15 H/cm2/s for 5000 s. Then TDS at 50 C/min to 1000 C.
See T. Hino et al., Fus. Engr. & Des. 39-40 (1998) pp.227-233.
end of title input
$ -----
main input
$ -----
dspcnme=h,end
sspcnme=h2,end
espcnme=h2g,end
segnds=12,18,end          $ 1 implant zone 15 nm, 2 bulk 0.1 mm
nbrencl=3,end             $ 1 test chamber, 2 source, 3 sink
linksegs=1,2,end
end of main input
$ -----
enclosure input
$ -----
start func,1,end          $ Test chamber where sample is
$ Enclosure 1 is the plasma chamber with pressure assumed negligible
etemp=tabl,1,end
esppres=h2g,1.0e-5,end
evol=0.12,end             $ Fit value m3
outflow=nbrflwp,1,qflow,const,8.33e-4,rencl,2,end
$
start bdry,2,end
$ Enclosure 2 is the sink for the vacuum pumping system
etemp=const,300.,end
esppres=h2g,const,1.0e-8,end
$
start bdry,3,end
$ Enclosure 3 is the source for the test chamber (ion source)
etemp=const,300.,end
esppres=h2g,const,1.0e-5,end
outflow=nbrflwp,1,qflow,const,8.33e-4,rencl,1,end
end of enclosure input
$ -----
thermal input
$ -----
start thermseg,end
$ 15-nm implantation zone [THERMSEG 1]
delx=0.0,10*1.5e-9,0.0,end
tempd=12*300.,end        $ Initial temperatures=(K)
tcon=equ,1,end           $ W thermal cond. (W/m-K)
rhocp=equ,2,end          $ rho*cp for W (J/m3K)
hsrc=const,0.,srcpf,12*0.,end $ Neglect internal heat sources
htrbcl=stemp,tabl,1,end  $ Temperature at the plasma-side surface
htrbcr=link,end
hgap=const,1.e9,end      $ Effectively infinite gap conductance
$
start thermseg,end
$ Balance of 0.1-mm tungsten specimen [THERMSEG 2]
delx=0.,1.e-9,1.e-8,1.0e-7,1.0e-6,12*7.407e-6,0.0,end
tempd=18*300.,end        $ Initial temperatures=(K)

```

```

tcon=equ,1,end          $ W thermal cond. (W/m-K)
rhocp=equ,2,end         $ rho*cp for W (J/m3K)
hsrcc=const,0.,srcpf,18*0.,end $ Neglect internal heat sources
htrbcl=link,end         $ Temperature at the plasma-side surface
htrbcr=stemp,tabl,1,end $ Temperature at the back-side surface
$
end of thermal input
$ =====
diffusion input
$ =====
start diffseg,end
$ 15-nm implantation zone [DIFFSEG 1]
nbrden=6.25e28,end
concd=h,const,1.0e-10,end $ Starting mobile concentration
ssconc=h2,1.0,link,end $ Starting surface species concentration
trapping=ttyp,1,tconc,const,0.07,tspc,h,alphr,equ,4
      alphr,equ,3,ctrap,const,0.9
      ttyp,2,tconc,const,1.38e-5,tspc,h,alphr,equ,5
      alphr,equ,3,ctrap,const,0.9
      ttyp,3,tconc,const,5.9e-6,tspc,h,alphr,equ,6
      alphr,equ,3,ctrap,const,0.99,end
qstrdr=h,const,0.,end $ Q*/R for Soret effect unknown
dcoef=h,equ,7,h2,equ,10,end $ Diffusion coeff (m2/s) [Modified]
srcsd=h,tabl,2,srcpf,norm,1.0,4.6e-9,3.0e-9,0.0,end
difbcl=surfdep,encl,1
      spc,h,nu,8.4e12,ec,-0.8,es,1.04
      comb,h,prob,1.0
      spc,h2,nu,8.4e12,ec,-0.1
      exch,h2g,amu,2.0,ex,0.05
      diss,h,h,eb,0.05
      form,h,h,prob,1.0,end
difbcr=link,h,solcon,equ,8,end
surfa=4.025e-5,end $ 14 mm diameter beam (derived)
$
start diffseg,end
$ Balance of 0.1-mm tungsten specimen [DIFFSEG 2]
nbrden=6.25e28,end
concd=h,const,1.0e-10,end $ Starting mobile concentration
ssconc=h2,link,1.0,end $ Starting surface species concentration
trapping=ttyp,1,tconc,const,1.38e-5,tspc,h,alphr,equ,5
      alphr,equ,3,ctrap,const,0.9
      ttyp,2,tconc,const,5.9e-6,tspc,h,alphr,equ,6
      alphr,equ,3,ctrap,const,0.99,end
qstrdr=h,const,0.,end $ Q*/R for Soret effect unknown
dcoef=h,equ,9,h2,equ,10,end $ Diffusion coeff (m2/s) [Modified]
srcsd=h,const,0.0,srcpf,18*0.0,end
difbcr=surfdep,encl,1
      spc,h,nu,8.4e12,ec,-0.8,es,1.04
      comb,h,prob,1.0
      spc,h2,nu,8.4e12,ec,-0.1
      exch,h2g,amu,2.0,ex,0.05
      diss,h,h,eb,0.05
      form,h,h,prob,1.0,end
difbcl=link,h,solcon,equ,8,end
surfa=4.025e-5,end $ 14 mm diameter beam (derived)
$
end of diffusion input

```

```

$
equation input
$ (1) Thermal conductivity of tungsten (W/m-K)
y=163.-0.0739*temp+2.89e-5*temp**2-4.3e-9*temp**3,end
$ (2) Rho Cp for tungsten (J/m3K)
y=(1930.-.0388*temp)*(131.+0.0226*temp-5.73e-6*temp**2+3.69e-9
*temp**3),end
$ (3) Alpht for h in tungsten (1/s)
y=9.1316e12*exp(-0.39/8.625e-5/temp),end
$ (4) Alphr for trap 1 in tungsten (1/s)
y=8.4e12*exp(-1.0/8.625e-5/temp),end
$ (5) Alphr for trap 2 in tungsten (1/s)
y=8.4e12*exp(-1.35/8.625e-5/temp),end
$ (6) Alphr for trap 3 in tungsten (1/s)
y=8.4e12*exp(-2.7/8.625e-5/temp),end
$ (7) Diffusivity for h in tungsten (m2/s)
y=4.1e-7*exp(-0.39/8.625e-5/temp),end
$ (8) Hydrogen solubility in tungsten (1/m3-Pa1/2)
y=1.83e24*exp(-1.04/8.625e-5/temp),end
$ (9) Diffusivity for h in implant-layer tungsten (m2/s)[Modified]
y=4.1e-7*exp(-.39/8.625e-5/temp)*10.,end
$ (10) Surface diffusivity for h2 at tungsten surface (m2/s)
y=4.1e-7*exp(-.1/8.625e-5/temp),end
end of equation input
$
table input
$ (1) Temperature history of enclosure 1
0.,300.,7500.,300.,8667.,1273.,1.0e10,1273.,end
$ (2) Implantation flux history (atom/m2/s)
0.,1.e19,5000.,1.e19,5001.,0.0,1.0e10,0.0,end
end of table input
$
control input
time=0.,end
tstep=1.0,end           $ time step = 1 sec
timend=9250.0,end       $ after implantation and desorption
nprint=120,end          $ print every 20 minutes
itermx=10000,end
delcmx=1.e-6,end
bump=1.e-4,end
bound=5.0,end
omega=0.9,end
damp=0.9,end
end of control input
$
plot input
nplot=10,end            $ makes plotfile entry every 10 sec
plotseg=1,2,end         $ segments for which plot info is needed
plotenc1=1,end          $ enclosures for which plot info is needed
dname=h,end             $ diffusing species for which plot info is needed
ename=h2g,end           $ enclosure species for which plot info is needed
dplot=moblinv,trapinv,sflux,stemp,end
eplot=press,diff,end     $ flow of molecules into enclosure not needed
end of plot input
$
end of data

```

Co-permeation of H and D through Pd (Val-2ea, see p. 53)

```
title input
  Sample Problem #5a - Co-Permeation of D and H throu Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.05 mm, 825 K, D2 only
end of title input
$
main input
  dspcnme=d,end
  espcnme=d2,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espPRES d2,1.0e-6,end
  outflow=nbrflwp,2,qflow,const,0.1,rencl,2
  qflow,const,0.1,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espPRES=d2,1.e-6,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espPRES=d2,1.e-6,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espPRES d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espPRES=d2,tabl,2,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*5.0e-6,0.0,end
  tempd=12*300.0,end
  tcon=const,73.,end
  rhocp=const,2.932e6,end
```

```

        hsrc=const,0.0,srcpf,12*0.0,end
        htrbcl=stemp,tabl,1,end
        htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
    nbrden=6.806e28,end
    concd=d,12*1.0e5,end
    dcoef=d,equ,2,end
    qstrdr=d,const,0.0,end
    srcsd=d,const,0.0,srcpf,12*0.0,end
    difbcl=lawdep,encl,2
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    difbcr=lawdep,encl,3
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of H,D in Pd
$(E. M. Wise, 1968, Palladium Recovery, Properties, and Uses, Academic Press,
$ New York, pp. 149-157.)
y=1.082e26*exp(-5000./temp),end
$
end of equation input
$
$
table input
$ (1) Temperature history
0.0,825.,8.e5,825.,end
$ (2) Pressure history of D2 in Enclosure 5
0.0,1.20e-04,150.,1.20e-4,151.,2.41e-4,250.,2.41e-4,251.,6.06e-4,350.,6.06e-4
351.,1.30e-3,450.,1.30e-3,451.,2.53e-3,550.,2.53e-3,551.,7.08e-3,650.
7.08e-3,651.,1.45e-2,750.,1.45e-2,751.,2.63e-2,850.,2.63e-2,851.,6.51e-2
950.,6.51e-2,951.,0.116,1050.,0.116,1051.,0.297,1150.,0.297,1151.,0.76,
1250.,0.76,1251.,1.55,1350.,1.55,1351.,3.37,1900.,3.37,end
end of table input
$
$
control input
    time=0.0,end
    tstep=0.1,end
    timend=1450.0,end
    nprint=500,end

```

```

    itermx=9000,end
    delcmx=1.0e-7,end
    bump=1.e-3,end
    damp=0.7
    bound=4.0,end
    omega=1.3,end
end of control input
$
$
plot input
    nplot=1000,end
    plotseg=1,end
    plotenc1=2,3,5,end
    dname=d,end
    ename=d2,end
    dplot=moblinv,sflux,end
    eplot=press,conv,diff,end
end of plot input
$
end of data

```


Co-permeation of H and D through Pd (Val-2eb, see p. 53)

```
title input
  Sample Problem #5b - Co-Permeation of D and H throu Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.025 mm, 825 K, D2 only
end of title input
$
main input
  dspcnme=d,end
  espcnme=d2,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espres d2,1.0e-6,end
  outflow=nbrflwp,2,qflow,const,0.1,rencl,2
  qflow,const,0.1,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espres=d2,1.e-6,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espres=d2,1.e-6,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espres d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espres=d2,tabl,2,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*2.5e-6,0.0,end
  tempd=12*300.0,end
  tcon=const,73.,end
```

```

        rhocp=const,2.932e6,end
        hsrc=const,0.0,srcpf,12*0.0,end
        htrbcl=stemp,tabl,1,end
        htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
    nbrden=6.806e28,end
    concd=d,12*1.0e5,end
    dcoef=d,equ,2,end
    qstrdr=d,const,0.0,end
    srcsd=d,const,0.0,srcpf,12*0.0,end
    difbcl=lawdep,encl,2
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    difbcr=lawdep,encl,3
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of H,D in Pd
$(E. M. Wise, 1968, Palladium Recovery, Properties, and Uses, Academic Press,
$ New York, pp. 149-157.)
y=1.082e26*exp(-5000./temp),end
$
end of equation input
$
$
table input
$ (1) Temperature history
0.0,825.,8.e5,825.,end
$ (2) Pressure history of D2 in Enclosure 5
0.0,1.00e-04,150.,1.00e-4,151.,2.37e-4,250.,2.37e-4,251.,5.71e-4,350.,5.71e-4
351.,1.24e-3,450.,1.24e-3,451.,2.53e-3,550.,2.53e-3,551.,6.87e-3,650.
6.87e-3,651.,.0128,750.,.0128,751.,2.63e-2,850.,2.63e-2,851.,6.61e-2
950.,6.61e-2,951.,0.118,1050.,0.118,1051.,0.302,1150.,0.302,1151.,0.76
1250.,0.76,1251.,1.55,1350.,1.55,1351.,3.37,1900.,3.37,end
$
end of table input
$
$
control input
    time=0.0,end
    tstep=0.1,end

```

```

    timend=1450.0,end
    nprint=500,end
    itermx=9000,end
    delcmx=1.0e-7,end
    bump=1.e-3,end
    damp=0.7
    bound=4.0,end
    omega=1.3,end
end of control input
$
$
plot input
    nplot=1000,end
    plotseg=1,end
    plotencl=2,3,5,end
    dname=d,end
    ename=d2,end
    dplot=moblinv,sflux,end
    eplot=press,conv,diff,end
end of plot input
$
end of data

```

Co-permeation of H and D through Pd (Val-2ec, see p. 53)

```
title input
  Sample Problem #5c - Co-Permeation of D and H through Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.025 mm, 865 K, D2 only
end of title input
$
main input
  dspcnme=d,end
  espcnme=d2,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espres d2,1.0e-6,end
  outflow=nbrflwp,2,qflow,const,0.1,rencl,2
  qflow,const,0.1,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espres=d2,1.e-6,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espres=d2,1.e-6,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espres d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espres=d2,tabl,2,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*2.5e-6,0.0,end
  tempd=12*300.0,end
  tcon=const,73.,end
```

```

        rhocp=const,2.932e6,end
        hsrc=const,0.0,srcpf,12*0.0,end
        htrbcl=stemp,tabl,1,end
        htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
    nbrden=6.806e28,end
    concd=d,12*1.0e5,end
    dcoef=d,equ,2,end
    qstrdr=d,const,0.0,end
    srcsd=d,const,0.0,srcpf,12*0.0,end
    difbcl=lawdep,encl,2
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    difbcr=lawdep,encl,3
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of H,D in Pd
$(E. M. Wise, 1968, Palladium Recovery, Properties, and Uses, Academic Press,
$ New York, pp. 149-157.)
y=1.082e26*exp(-5000./temp),end
$
end of equation input
$
$
table input
$ (1) Temperature history
0.0,865.,8.e5,865.,end
$ (2) Pressure history of D2 in Enclosure 5
0.0,1.00e-04,150.,1.00e-4,151.,2.37e-4,250.,2.37e-4,251.,5.71e-4,350.,5.71e-4
351.,1.24e-3,450.,1.24e-3,451.,2.53e-3,550.,2.53e-3,551.,6.87e-3,650.
6.87e-3,651.,.0128,750.,.0128,751.,2.63e-2,850.,2.63e-2,851.,6.61e-2
950.,6.61e-2,951.,0.118,1050.,0.118,1051.,0.302,1150.,0.302,1151.,0.76
1250.,0.76,1251.,1.55,1350.,1.55,1351.,3.37,1900.,3.37,end
$
end of table input
$
$
control input
    time=0.0,end
    tstep=0.1,end

```

```

    timend=1450.0,end
    nprint=500,end
    itermx=9000,end
    delcmx=1.0e-7,end
    bump=1.e-3,end
    damp=0.7
    bound=4.0,end
    omega=1.3,end
end of control input
$
$
plot input
    nplot=1000,end
    plotseg=1,end
    plotencl=2,3,5,end
    dname=d,end
    ename=d2,end
    dplot=moblinv,sflux,end
    eplot=press,conv,diff,end
end of plot input
$
end of data

```

Co-permeation of H and D through Pd (Val-2ed, see p. 53)

```
title input
  Sample Problem #5d - Co-Permeation of D and H through Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.025 mm, 870 K, H2, D2, and HD present, lawdep diffusion bc.
end of title input
$
main input
  dspcnme=h,d,end
  espcnme=h2,d2,hd,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espPRES h2,1.0e-7,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,2,qflow,const,0.1,rencl,2
  qflow,const,0.1,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espPRES h2,0.063,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
  espcomb=hd,const,2.0,h2,0.5,d2,0.5,end
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espPRES h2,1.0e-7,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
  espcomb=hd,const,2.0,h2,0.5,d2,0.5,end
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espPRES h2,1.e-10,hd,1.e-10,d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espPRES=h2,tabl,3,hd,const,1.0e-10,d2,tabl,2,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*2.5e-6,0.0,end
```

```

tempd=12*300.0,end
tcon=const,73.,end
rhocp=const,2.932e6,end
hsrcc=const,0.0,srcpf,12*0.0,end
htrbcl=stemp,tabl,1,end
htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
  nbrden=6.806e28,end
  concd=h,12*0.0,d,12*0.0,end
  dcoef=h,equ,1,d,equ,2,end
  qstrdr=h,const,0.0,d,const,0.0,end
  srccd=h,const,0.0,srcpf,12*0.0,d,const,0.0,srcpf,12*0.0,end
  difbcl=lawdep,encl,2
    dspc,h,h2,pexp,0.8958,solcon,equ,3
    dspc,d,d2,pexp,0.8958,solcon,equ,3,end
  difbcr=lawdep,encl,3
    dspc,h,h2,pexp,0.8958,solcon,equ,3
    dspc,d,d2,pexp,0.8958,solcon,equ,3,end
  surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of H/D in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp),end
$
end of equation input
$
$
table input
$ (1) Temperature history
0.0,870.,8.e5,870.,end
$ (2) Pressure history of D2 in Enclosure 5
0.,1.8421e-4,150.,1.8421e-4,151.,1.37e-3,250.,1.37e-3,251.,4.8193e-3,350.
4.8193e-3,351.,0.022124,450.,0.022124,451.,0.06135,550.,0.06135,551.
0.15209,650.,0.15209,651.,0.44405,750.,0.44405,751.,0.94073,1.e6,0.94073,end
$ (3) Pressure history of H2 in Enclosure 5
0.,0.05968,150.,0.05968,151.,0.05437,250.,0.05437,251.,0.04782,350.
0.04782,351.,0.03512,450.,0.03512,451.,0.02435,550.,0.02435,551.
0.01509,650.,0.01509,651.,0.00705,750.,0.00705,751.,0.003734,1.e6,0.003734,en
d
end of table input
$

```



```

$
control input
  time=0.0,end
  timestep=0.1,end
  timend=801.0,end
  nprint=1000,end
  itermx=9000,end
  delcmx=1.0e-6,end
  bump=1.e-3,end
  damp=0.2
  bound=9.0,end
  omega=0.3,end
end of control input
$
$
plot input
  nplot=1000,end
  plotseg=1,end
  plotencl=2,3,5,end
  dname=h,d,end
  ename=h2,hd,d2,end
  dplot=moblinv,sflux,end
  eplot=press,conv,diff,end
end of plot input
$
end of data

```

Co-permeation of H and D through Pd (Val-2eea, see p. 53)

```
title input
  Sample Problem #5e - Co-Permeation of D and H through Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.025 mm, 870 K, H2, D2, and HD present, ratedep diffusion bc
end of title input
$
main input
  dspcnme=h,d,end
  espcnme=h2,d2,hd,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espPRES h2,1.0,hd,1.0,d2,1.0,end
  outflow=nrflwp,2,qflow,const,1.e-8,rencl,2
  qflow,const,1.e-8,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espPRES h2,0.063,hd,2.0e-7,d2,1.0e-7,end
  outflow=nrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espPRES h2,1.0e-7,hd,2.0e-7,d2,1.0e-7,end
  outflow=nrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espPRES h2,1.e-10,hd,1.e-10,d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espPRES=h2,const,0.063,hd,const,1.0e-10,d2,tabl,2,end
  outflow=nrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*2.5e-6,0.0,end
  tempd=12*300.0,end
  tcon=const,73.,end
```

```

        rhocp=const,2.932e6,end
        hsrc=const,0.0,srcpf,12*0.0,end
        htrbcl=stemp,tabl,1,end
        htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
    nbrden=6.806e28,end
    concd=h,12*1.0,d,12*1.0,end
    dcoef=h,equ,1,d,equ,2,end
    qstrdr=h,const,0.0,d,const,0.0,end
    srcsd=h,const,0.0,srcpf,12*0.0,d,const,0.0,srcpf,12*0.0,end
    difbcl=ratedep,encl,2
        spc,h
            exch,h2,ksubd,equ,4
                h,ksubr,equ,7
            exch,hd,ksubd,equ,5
                d,ksubr,equ,9
        spc,d
            exch,d2,ksubd,equ,6
                d,ksubr,equ,8
            exch,hd,ksubd,equ,5
                h,ksubr,equ,9,end
    difbcr=ratedep,encl,3
        spc,h
            exch,h2,ksubd,equ,4
                h,ksubr,equ,7
            exch,hd,ksubd,equ,5
                d,ksubr,equ,9
        spc,d
            exch,d2,ksubd,equ,6
                d,ksubr,equ,8
            exch,hd,ksubd,equ,5
                h,ksubr,equ,9,end
    surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of D in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp),end
$
$ (4) Dissociation coefficient for H2
y=2.6276e24/sqrt(2.*temp),end
$

```

```

$ (5) Dissociation coefficient for HD
y=2.6276e24/sqrt(3.*temp),end
$
$ (6) Dissociation coefficient for D2
y=2.6276e24/sqrt(4.*temp),end
$
$ (7) Recombination coefficient H2
y=2.2444e-28/sqrt(2.*temp)*exp(10000./temp)*1.23**2,end
$
$ (8) Recombination coefficient D2
y=2.2444e-28/sqrt(4.*temp)*exp(10000./temp),end
$
$ (9) Recombination coefficient HD
y=2.2444e-28/sqrt(3.*temp)*exp(10000./temp)*1.125**2,end
$
$ (10) Solubility of h in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp)*1.23,end
$
end of equation input
$
$
table input
$ (1) Temperature history
0.0,870.,8.e5,870.,end
$ (2) Pressure history of D2 in Enclosure 5
0.,0.0035,150.,0.0035,151.,0.01,250.,0.01,251.,0.02,350.,0.02,351.
0.05,450.,0.05,451.,0.1,550.,0.1,551.,0.2,650.,0.2,651.,0.5,750.,0.5
751.,0.9,1.e6,0.9,end
end of table input
$
$
control input
    time=0.0,end
    tstep=0.1,end
    timend=900.0,end
    nprint=1000,end
    itermx=9000,end
    delcmx=1.0e-6,end
    bump=1.e-3,end
    damp=0.2
    bound=4.0,end
    omega=0.3,end
end of control input
$
$
plot input
    nplot=1000,end
    plotseg=1,end
    plotencl=2,3,5,end
    dname=h,d,end
    ename=h2,hd,d2,end
    dplot=moblinv,sflux,sconc,end
    eplot=press,conv,diff,end
end of plot input
$
end of data

```

Co-permeation of H and D through Pd (Val-2eeb, see p. 53)

```
title input
  Sample Problem #5e - Co-Permeation of D and H throu Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.025 mm, 870 K, H2, D2, and HD present, ratedep diffusion bc
end of title input
$
main input
  dspcnme=h,d,end
  espcnme=h2,d2,hd,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espPRES h2,1.0,hd,1.0,d2,1.0,end
  outflow=nbrflwp,2,qflow,const,1.e-8,rencl,2
  qflow,const,1.e-8,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espPRES h2,0.063,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espPRES h2,1.0e-7,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espPRES h2,1.e-10,hd,1.e-10,d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espPRES=h2,const,0.063,hd,const,1.0e-10,d2,tabl,2,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*2.5e-6,0.0,end
  tempd=12*300.0,end
  tcon=const,73.,end
  rhocp=const,2.932e6,end
```

```

        hsrc=const,0.0,srcpf,12*0.0,end
        htrbcl=stemp,tabl,1,end
        htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
    nbrden=6.806e28,end
    concd=h,12*1.0,d,12*1.0,end
    dcoef=h,equ,1,d,equ,2,end
    qstrdr=h,const,0.0,d,const,0.0,end
    srcsd=h,const,0.0,srcpf,12*0.0,d,const,0.0,srcpf,12*0.0,end
    difbcl=ratedep,encl,2
        spc,h
            exch,h2,ksubd,equ,4
                h,ksubr,equ,7
            exch,hd,ksubd,equ,5
                d,ksubr,equ,9
        spc,d
            exch,d2,ksubd,equ,6
                d,ksubr,equ,8
            exch,hd,ksubd,equ,5
                h,ksubr,equ,9,end
    difbcr=ratedep,encl,3
        spc,h
            exch,h2,ksubd,equ,4
                h,ksubr,equ,7
            exch,hd,ksubd,equ,5
                d,ksubr,equ,9
        spc,d
            exch,d2,ksubd,equ,6
                d,ksubr,equ,8
            exch,hd,ksubd,equ,5
                h,ksubr,equ,9,end
    surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of D in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp),end
$
$ (4) Dissociation coefficient for H2
y=2.6276e24/sqrt(2.*temp)/10.,end
$
$ (5) Dissociation coefficient for HD

```

```

y=2.6276e24/sqrt(3.*temp)/10.,end
$
$ (6) Dissociation coefficient for D2
y=2.6276e24/sqrt(4.*temp)/10.,end
$
$ (7) Recombination coefficient H2
y=2.2444e-28/sqrt(2.*temp)*exp(10000./temp)/10.*1.23**2,end
$
$ (8) Recombination coefficient D2
y=2.2444e-28/sqrt(4.*temp)*exp(10000./temp)/10.,end
$
$ (9) Recombination coefficient HD
y=2.2444e-28/sqrt(3.*temp)*exp(10000./temp)/10.*1.125**2,end
$
$ (10) Solubility of H in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp)*1.23,end
$
end of equation input
$
$
table input
$ (1) Temperature history
0.0,870.,8.e5,870.,end
$ (2) Pressure history of D2 in Enclosure 5
0.,0.0035,150.,0.0035,151.,0.01,250.,0.01,251.,0.02,350.,0.02,351.
0.05,450.,0.05,451.,0.1,550.,0.1,551.,0.2,650.,0.2,651.,0.5,750.,0.5
751.,0.9,1.e6,0.9,end
end of table input
$
$
control input
    time=0.0,end
    tstep=0.1,end
    timend=900.0,end
    nprint=1000,end
    itermx=9000,end
    delcmx=1.0e-6,end
    bump=1.e-3,end
    damp=0.2
    bound=4.0,end
    omega=0.3,end
end of control input
$
$
plot input
    nplot=1000,end
    plotseg=1,end
    plotencl=2,3,5,end
    dname=h,d,end
    ename=h2,hd,d2,end
    dplot=moblinv,sflux,sconc,end
    eplot=press,conv,diff,end
end of plot input
$
end of data

```

Co-permeation of H and D through Pd (Val-2ef, see p. 53)

```
title input
  Sample Problem #5e - Co-Permeation of D and H throu Pd by K. Kizu,
  A. Pisarev and T. Tanabe, Journal of Nuclear Materials, 289 (2001) 291-302.
  Pd 0.025 mm, 870 K, H2, D2, and HD present, mixed lawdep(up)/ratedep (down)
diffusion bc
end of title input
$
main input
  dspcnme=h,d,end
  espcnme=h2,d2,hd,end
  segnds=12,end
  nbrencl=5,end
end of main input
$
enclosure input
$
start bdry,1,end
$ This is the background pressure source for both active chambers
  etemp=tabl,1,end
  espPRES h2,1.0,hd,2.0,d2,1.0,end
  outflow=nbrflwp,2,qflow,const,1.0e-8,rencl,2
  qflow,const,1.0e-8,rencl,3,end
$
start func,2,end
$ This is the upstream chamber connecting to the membrane
  etemp=tabl,1,end
  espPRES h2,0.0063,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
  espcomb=hd,const,2.0,h2,0.5,d2,0.5,end
$
start func,3,end
$ This is the downstream chamber connected to the membrane
  etemp=tabl,1,end
  espPRES h2,1.0e-7,hd,2.0e-7,d2,1.0e-7,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,4,end
  evol=0.05,end          $ Estimated volume
$
start bdry,4,end
$ This is the gas sink representing the vacuum pumping system
  etemp=tabl,1,end
  espPRES h2,1.e-10,hd,1.e-10,d2,1.e-10,end
$
start bdry,5,end
$ This is the gas source with pre-programmed species pressures
  etemp=tabl,1,end
  espPRES=h2,tabl,3,hd,const,1.0e-10,d2,tabl,2,end
  outflow=nbrflwp,1,qflow,const,0.1,rencl,2,end
end of enclosure input
$
thermal input
$ Segment 1 - Pd film
start thermseg,end
  delx=0.0,10*2.5e-6,0.0,end
  tempd=12*300.0,end
```



```

        tcon=const,73.,end
        rhocp=const,2.932e6,end
        hsrc=const,0.0,srcpf,12*0.0,end
        htrbcl=stemp,tabl,1,end
        htrbcr=stemp,tabl,1,end
end of thermal input
$
$
diffusion input
$ Segment 1 - Pd flim
start diffseg,end
    nbrden=6.806e28,end
    concd=h,12*1.0,d,12*1.0,end
    dcoef=h,equ,1,d,equ,2,end
    qstrdr=h,const,0.0,d,const,0.0,end
    srcsd=h,const,0.0,srcpf,12*0.0,d,const,0.0,srcpf,12*0.0,end
    difbcl=lawdep,encl,2
        dspc,h,h2,pexp,0.8958,solcon,equ,10
        dspc,d,d2,pexp,0.8958,solcon,equ,3,end
    difbcr=ratedep,encl,3
        spc,h
            exch,h2,ksubd,equ,4
            h,ksubr,equ,7
            exch,hd,ksubd,equ,5
            d,ksubr,equ,9
        spc,d
            exch,d2,ksubd,equ,6
            d,ksubr,equ,8
            exch,hd,ksubd,equ,5
            h,ksubr,equ,9,end
    surfa=1.8e-4,end
end of diffusion input
$
$
equation input
$
$ (1) Diffusivity of H in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960)]
y=4.31e-7*exp(-2818./temp),end
$
$ (2) Diffusivity of D in Pd [O. M. Katz & E. A. Gulbransen,Rev. Sci. Instr.,
$ 31, 615-617 (1960); divided by 1.414 for isotope effect]
y=3.048e-7*exp(-2818./temp),end
$
$ (3) Solubility of D in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp),end
$
$ (4) Dissociation coefficient for H2
y=2.6276e24/sqrt(2.*temp)/2.,end
$
$ (5) Dissociation coefficient for HD
y=2.6276e24/sqrt(3.*temp)/2.,end
$
$ (6) Dissociation coefficient for D2
y=2.6276e24/sqrt(4.*temp)/2.,end
$
$ (7) Recombination coefficient H2

```

```

y=2.2444e-28/sqrt(2.*temp)*exp(10000./temp)/2.*1.23**2,end
$
$ (8) Recombination coefficient D2
y=2.2444e-28/sqrt(4.*temp)*exp(10000./temp)/2.,end
$
$ (9) Recombination coefficient HD
y=2.2444e-28/sqrt(3.*temp)*exp(10000./temp)/2.*1.125**2,end
$
$ (10) Solubility of H in Pd based on measurements of Kizu et al.
y=1.082e26*exp(-5000./temp)*1.23,end
$
end of equation input
$
table input
$ (1) Temperature history
0.0,870.,8.e5,870.,end
$ (2) Pressure history of D2 in Enclosure 5
0.,1.8421e-4,150.,1.8421e-4,151.,1.37e-3,250.,1.37e-3,251.,4.8193e-3,350.
4.8193e-3,351.,0.022124,450.,0.022124,451.,0.06135,550.,0.06135,551.
0.15209,650.,0.15209,651.,0.44405,750.,0.44405,751.,0.94073,1.e6,0.94073,end
$ (3) Pressure history of H2 in Enclosure 5
0.,0.05968,150.,0.05968,151.,0.05437,250.,0.05437,251.,0.04782,350.
0.04782,351.,0.03512,450.,0.03512,451.,0.02435,550.,0.02435,551.
0.01509,650.,0.01509,651.,0.00705,750.,0.00705,751.,0.003734,1.e6,0.003734,en
d
end of table input
$
$
control input
time=0.0,end
tstep=0.1,end
timend=900.0,end
nprint=1000,end
itermx=1000,end
delcmx=1.0e-6,end
bump=1.e-3,end
damp=0.1
bound=4.0,end
omega=0.1,end
end of control input
$
$
plot input
nplot=1000,end
plotseg=1,end
plotencl=2,3,5,end
dname=h,d,end
ename=h2,hd,d2,end
dplot=moblinv,sflux,sconc,end
eplot=press,conv,diff,end
end of plot input
$
end of data

```